Turkish Society of Hematology
1st International Lymphoma - Leukemia - Myeloma (LLM) Congress
Proceedings and Abstract Book
May 24 - 27, 2007
Lykia World - Ölüdeniz, Fethiye / Turkey
TURKISH SOCIETY OF HEMATOLOGY
40TH ANNIVERSARY

1ST INTERNATIONAL
LYMPHOMA - LEUKEMIA - MYELOMA (LLM) CONGRESS

Proceedings and Abstract Book

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Lykia World - Ölüdeniz, Fethiye / Turkey
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Dear Colleagues,

It is our pleasure to welcome you to 1st International Congress of Leukemia-Lymphoma-Myeloma in Fethiye, Turkey.

We are so proud to announce that the proceedings and the abstracts of the 1st International Congress of Leukemia-Lymphoma-Myeloma are published in the Turkish Journal of Hematology.

“Turkish Journal of Hematology” plays an important role in the field of hematology by means of publishing sections of research and review articles, case reports and images in hematology. The supplement book of Turkish Journal Hematology comprises the work of the both speakers and the participants of the congress.

We believe that the congress have an outstanding scientific program. We wish to leave an important post congress source for the young researchers to use as a guide for his research life. This supplement of this congress has a higher priority for us, in which we believe this issue will serve as a perfect educational source for young fellows all around the world and also for us.

With the support of speakers, chairs, companies and abstract presenters the majority of the original program has been maintained and we are grateful to everyone for the strong commitment shown to the success of the meeting.

I would like to express my gratitude to all contributors. We hope that having this supplementary issue of the Turkish Journal Hematology is going to be useful for both you and your colleagues in the diagnosis and the treatment of the patients with hematological disorders.

Aytemiz Gürgey, MD
Editor in Chief
Turkish Journal of Hematology
May 24th, 2007, Thursday

HALL A
08:30 – 10:30  **Acute Myeloblastic Leukemia**
  **Chairs: John Kersey, Ali Ünal**
  Recent Advances in the Biology of AML – John Kersey, USA
  Chemotherapy in AML – Peter H. Wiernik, USA
  Autologous Stem Cell Transplantation in AML – Norbert Claude Gorin, France
  Allogeneic Stem Cell Transplantation in AML – Francesco Frassoni, Italy

SCIENTIFIC SUBCOMMITTEE MEETING
10:00 – 12:30  Myeloma Scientific Subcommittee Meeting

10:30 – 10:45  *Break*

HALL A
10:45 – 12:15  **Acute Lymphoblastic Leukemia**
  **Chairs: Oliver Ottman, İsmet Aydoğan**
  Ph⁺ ALL – Oliver Ottman, Germany
  Treatment of Adult ALL – Charles Linker, USA
  Treatment of Pediatric ALL – Giorgio Dini, Italy

12:15 – 13:15  *Lunch*

HALL B
13:15 – 14:15  **Satellite Symposia, Janssen Cilag**
  **Optimizing treatment strategies in Multiple Myeloma**
  New Agents in Front Line Multiple Myeloma – Jean Luc Harousseau, France
  Targeted Therapy and Tailored Treatment with Proteasome Inhibition in Multiple Myeloma – Orhan Sezer, Germany
  Maximizing the Benefits of Bortezomib at First Relaps Multiple Myeloma – Ali Ünal, Turkey

14:15 – 14:30  *Break*

HALL B
14:30 – 15:30  **Satellite Symposia, Bristol Myers Squibb**
  **New developments for patients with resistant Ph⁺ CML and Ph⁺ ALL**
  **Chair: Osman İlhan, Turkey**
  The role of new TKIs in imatinib resistant patients and imatinib intolerant patients Jane Apperley, UK
  Optimizing treatment decisions in treating Ph⁺ ALL – Zafer Gülbaş, Turkey
15:30 – 15:45  Break

**SCIENTIFIC SUBCOMMITTEE MEETING**

15:30 – 17:30  Chronic Leukemia Scientific Subcommittee Meeting

**HALL A**

15:45 – 17:15  Myeloma - I
   **Chairs: Jean Luc Harousseau, Meral Beksaç**
   Myeloma Biology and Molecular Pathology – Mohamad Mohty, France
   Bone Diseases and Treatment – Orhan Sezer, Germany
   Relapse and Refractory Myeloma – Jean Luc Harousseau, France

**HALL B**

15:45 – 17:15  Palliative Care / Supportive Therapies
   **Chairs: Claudio Viscoli, Hakan Özdoğu**
   Current Use of Erythropoietins in Hematological Malignancies – Pellegrino Musto, Italy
   The Role of Mucositis as a Predisposing Factor to Systemic Infection and Bacteremia – Claudio Viscoli, Italy
   Febrile Neutropenia in 2007 – Murat Akova, Turkey

**HALL A**

17:15 – 18:15  Interactive Case Discussions
   **Chair: Zafer Gülbaş**
   Acute Lymphoblastic Leukemia – Fahir Özkalemkaş, Turkey
   **Chair: Nejat Akar**
   Pediatric Acute Lymphoblastic Leukemia – Tiraje Celkan, Turkey

**HALL B**

17:15 – 18:15  Interactive Case Discussions
   **Chair: Serdar Bedii Omay**
   Follicular Lymphoma – Berksoy Şahin, Turkey
   **Chair: Mustafa Çetiner**
   Multiple Myeloma – Meral Beksaç, Turkey

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May 25th, 2007, Friday

**HALL A**

09:00 – 10:30  Follicular Lymphoma
   **Chairs: Anthony Goldstone, Bülent Ündar**
   Follicular Lymphoma – Pathology – İçiş Buluş, Turkey
   Treatment of Follicular Lymphoma – Eva Kimby, Sweden
   Follicular Lymphoma – High Dose Therapy – Anthony H. Goldstone, UK

10:30 – 10:45  Break

**SCIENTIFIC SUBCOMMITTEE MEETING**

10:00 – 12:30  Acut Leukemia Scientific Subcommittee Meeting

**HALL A**

10:45 – 12:15  Glressive Lymphoma
   **Chairs: Andreas Rosenwald, Semra Paydaş**
   Aggressive Lymphoma – Pathology – Andreas Rosenwald, Germany
   DLBCL-L First Line Treatment – Burhan Ferhanoğlu, Turkey
   DLBCL-L Relapse, Resistant Cases and Transplantation – Koen Van Besien, USA

12:15 – 13:15  Lunch


HALL B
13:15 – 14:15  **Satellite Symposia, Roche**
New Perspectives in Diffuse Large B Cell Lymphoma
Chair: Burhan Ferhanoğlu, Turkey
Michael Pfleunndschuh, Germany
14:15 – 14:30  **Break**

HALL A
14:30 – 16:00  **Hodgkin’s Disease**
*Chairs: Andreas Josting, Nilgün Sayinalp*
Hodgkin's Diseases First Line Treatment – Volker Diehl, Germany
Relapse and Refractory Hodgkin’s Diseases – Andreas Josting, Germany
16:00 – 16:15  **Break**

HALL A
16:15 – 17:15  **Myeloproliferative Disorders**
*Chairs: Yücel Tangün, Rauf Haznedar*
Biology Diagnosis and Classification of MPD – Johannes Jacobus Michiels, Belgium
The Targets of Therapy in Polycythemia Vera and Thrombocytemia – Tiziano Barbui, Italy

HALL A
17:15 – 18:15  **Interactive Case Discussions**
*Chair: Ayşen Timurağaoğlu*
Chronic Myeloid Leukemia – İbrahim Haznedaroğlu, Turkey
– Nilgün Sayinalp, Turkey
*Chair: Mehmet Ali Özcan*
Chronic Lymphocytic Leukemia – Bülent Ündar, Turkey

HALL B
17:15 – 18:15  **Interactive Case Discussions**
*Chair: Tanju Atamer*
Mantle Cell Lymphoma – Gülsan Sucak Türköz, Turkey
*Chair: Önder Arslan*
DLBCL – Mustafa Çetin, Turkey

May 26th, 2007, Saturday

HALL A
09:00 – 10:30  **Chronic Myeloid Leukemia**
*Chairs: Jane Apperley, İbrahim Haznedaroğlu*
CML: Case Closed? – Junia V. Melo, UK
Stem Cell Transplantation in CML – Jane Apperley, UK
Hype or Hope; Novel Tyrosine Kinease Inhibitors – Justus Duyster, Germany

**SCIENTIFIC SUBCOMMITTEE MEETING**
10:00 – 12:30  **Lymphoma Scientific Subcommittee Meeting**
10:30 – 10:45  **Break**
**HALL A**

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- **Tiziano Barbui, Zafer Gülbaş**
- **Levent Ündar, Turkey**
- **Sante Tura, Muhit Özcan**

- The Molecular Pathogenesis of MDS – Thomas Look, USA
- Current Non Transplant Treatment Strategies in MDS – Azra Raza, USA
- The Role of Allogeneic Stem Cell Transplantation in MDS – Theo de Witte, The Netherlands
- Optimal Bisphosphonate Treatment in Multiple Myeloma
- Practical Guidelines of the Therapy of CLL – Sante Tura, Italy
- Reduced-Intensity Conditioning Allogeneic Stem Cell Transplantation for LLM: Hype, Reality or Time for a Rethink? – Arnon Nagler, Israel
- Current Management of Hairy Cell Leukemia – Tadeusz Robak, Poland
- First Line Treatment of Multiple Myeloma – Guido Tricot, USA
- Stem Cell Transplantation in Multiple Myeloma-The German Experience – Herman Einsele, Germany

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**Chairs:**
- **Orhan Sezer, Levent Ündar**

- Current Status of Thalidomide in the Treatment of Multiple Myeloma – Mario Boccadoro, Italy
- Recent Advances of Thalidomide in the Treatment of Multiple Myeloma – Jean Luc Harousseau, France
- Thalidomide Treatment in Relapsed, Refractory Multiple Myeloma – Levent Ündar, Turkey
- First Line Treatment of Multiple Myeloma – Guido Tricot, USA
- Stem Cell Transplantation in Multiple Myeloma-The German Experience – Herman Einsele, Germany
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Chemotherapy in AML

Peter H. Wiernik
Our Lady of Mercy Cancer Center, New York Medical College, Bronx, New York, USA

Currently the best standard therapy for adults < 70 years of age consists of induction therapy with three daily doses of idarubicin and a seven-day continuous infusion of cytarabine. Some physicians still prefer daunorubicin or mitoxantrone instead of idarubicin, but all relevant prospective, randomized trials demonstrate one or more advantages of idarubicin over daunorubicin, and no studies demonstrate an advantage for mitoxantrone over daunorubicin. Furthermore, a meta-analysis of relevant raw data performed by Wheatley et al confirmed the superiority of idarubicin over daunorubicin as an induction agent. Most studies in which daunorubicin was used during induction employed three consecutive daily doses of 45 mg/M2. There is no evidence that 60 mg/M2 doses, as used by some investigators, lead to a better outcome than the lower doses. The standard dose of cytarabine used in induction is 100 mg/M2 daily, given as a continuous seven-day infusion. Doubling that dose, or even increasing it by a factor of 20 or 304 has resulted in little improvement, if any, in outcome of induction therapy. The addition of etoposide to the standard anthracycline + cytarabine induction regimen has improved results in some 5 but not all 6 studies.

There is general agreement that post-remission therapy is necessary to maximize disease-free and overall survival, but there is no universally accepted post-remission therapy regimen. High-dose cytarabine regimens have commonly been employed and seem to be effective, especially in younger patients with favorable cytogenetics. There is little evidence that combining other drugs with high-dose cytarabine post-remission improves results. The optimum dose of cytarabine as post-remission therapy has not been defined. It seems clear from the original study by Mayer et al that a dose of 400 mg/M2 is inferior to 3 gm/M2 but doses in between those have not been widely tested in an evaluable manner.

Despite the popularity of stem cell transplantation as a post-remission therapy, outcome data are disappointing for both autologous and allogeneic stem cell transplantation. In fact, Visani et al after an analysis of 344 papers concluded that there is no evidence that autologous stem cell transplantation is superior in terms of overall survival to chemotherapy alone, and that no overall benefit of allografting on survival was demonstrated by any trial. Also of note is the discovery that Hispanics allo-transplanted in the United States had a significantly higher risk of treatment failure (death or relapse) and overall mortality than Whites, for unknown reasons.

G-CSF and GM-CSF have both been shown not to worsen disease outcome when used as supportive care in patients with AML. On the other hand, they may have the potential for inducing secondary AML or myelodysplasia in certain solid tumor patients. A doubling of the incidence of AML/MDS in 5,510 women treated with adjuvant chemotherapy for breast cancer was observed in those who received colony-stimulating factors compared with those who did not.

Patients with AML over age 65 years generally have a poorer outcome with therapy than do
younger patients, and controversy exists as to whether older patients should be treated with regimens used in younger patients, or with less intensive therapy such as low-dose cytarabine. Kantarjian et al15 analyzed the data for 998 patients aged 65 years or more with AML or high-risk myelodysplasia treated with intensive therapy in an effort to determine prognostic factors for response and survival. The overall complete response rate was 45%. Poor prognostic factors for complete response and survival were age >75 years, unfavorable karyotype, poor performance status, longer duration of antecedent hematologic disorder and abnormal organ function. Based on these prognostic factors, they estimated that approximately 20% of the patients fell into a good prognosis group with an expected complete response rate > 60%, an induction mortality rate of 10% and a 1-year survival rate >50%. Such patients would clearly be expected to benefit from standard intensive therapy. Appelbaum et al16 studied a similar group of almost identical size. In addition to the prognostic factors noted above, they found multidrug resistance protein in 33% of AML patients < age 56 compared with 57% of patients older than 75 years. Consistent with the Kantarjian et al study15 they observed that 35% of patients younger than age 56 had unfavorable cytogenetics, compared with 51% of patients older than 75 years. It seems advisable to treat elderly AML patients with good prognostic factors as described in these two studies with standard induction chemotherapy. It is not as clear how to approach post-remission therapy. Standard high-dose cytarabine is too toxic for most elderly patients. Doses of 1.0-1.5 gm/M<sup>2</sup> have been well tolerated but not clearly effective.13

The best hope for improving therapy for adult AML is the development of new drugs with better activity against the disease. After a long drought, a number of recently introduced agents have already demonstrated promise. Giles et al17 studied cloretazine in patients age >60 years with previously untreated AML. The drug was given alone at a dose of 600 mg/M<sup>2</sup> once, as induction therapy to 104 patients with a median age of 72 years. No patient had a favorable karyotype, and most had some significant organ dysfunction. The complete response rate was 28% and another 4% had a complete response with incomplete recovery. The one-year survival rate for the 32% of patients who were complete responders was 28%. There was minimal extramedullary toxicity in the study. The drug causes DNA crosslinks. Its active metabolite has similarities to that of carmustine (BCNU) but it yields more than twice the DNA crosslinks, mole for mole, compared with carmustine.18 Burnett et al19 administered clofarabine (a purine nucleoside analog) 30 mg/M<sup>2</sup> daily for 5 days to 66 patients with a median age of 71 years. 62 had intermediate or poor risk cytogenetics. One course of drug was given every 28-42 days and a maximum of 3 courses were given. The CR + CRi rate was 29% and the one-year overall survival rate for responders was 32% and 28% for non-responders. Interesting, the one-year survival rate was identical for intermediate and poor cytogenetics patients. Clofarabine appears to be more toxic than cloretazine in the doses and schedules used. Serious renal toxicity developed in about 18% of patients treated with the former, and sepsis occurred in approximately 26% of those patients.

Several recent studies, if confirmed, will result in improved treatment of patients with AML in the near future. Liu et al20 assessed response and survival in 60 patients with APL induced with ATRA, 25 mg/M<sup>2</sup> plus As<sub>2</sub>O<sub>3</sub>, 0.16 mg/kg and consolidated them with 3 cycles of daunorubicin, cytarabine and homoharringtonine, and compared results with 56 historical controls induced with ATRA alone followed by postremission chemotherapy. The experimental group also received 5 cycles of maintenance therapy with monthly ATRA, followed by As<sub>2</sub>O<sub>3</sub> daily for a month, which was followed by weekly methotrexate for a month. There was no difference in CR rate between the groups, which was low (56% v 51%). However, at a median follow-up of 48 and 56 months, overall and event-free survival were significantly longer in the study group (4-year overall survival 98.1% v 83.4%, and 4-year event-free survival 94.2% v 45.6%).

The MRC21 studied the addition of gemtuzumab ozogamicin (GO), 3 mg/M<sup>2</sup> on day 1 of induction therapy with ADE, DA or FLAG-Ida in a randomized study of 113 patients <60 years old. CR rates were not different (85%). At 3 years, disease-free survival was significantly different in favor of those who received GO (49% v 38%). Toxicity was similar between the groups. Others22 have shown in vitro that cytotoxic activity of GO correlates with expression of protein kinase Syk and that azacytidine upregulates Syk. In another in vitro study Takahashi et al23 demonstrated a synergistic effect of As<sub>2</sub>O<sub>3</sub> and FLT 3 inhibition on cells with FLT 3-ITD.

Schlenk et al24 performed a retrospective analysis of 4 German AML Study Group trials. The studies were of similar design and included 872
patients with a median age of 48 years. The results of gene analyses indicated that the 33% of patients found to be NPM1+ and FLT3 ITD – as well as those CEBPA+ had significantly higher response rates than others (88% and 83% for the former and 66% for others). Furthermore, those favorable genotypes were associated with significantly better relapse-free and overall survival. Others 25 have confirmed in a larger study that if not associated with FLT3-ITD mutations, mutant NPM1 appears to identify patients with improved response to treatment.

References


A utologous hematopoietic stem cell transplantation remains presently an interesting therapeutic option in adult patients with AML beyond 35 years of age or if younger with no identical sibling for an allogeneic transplantation.

Data from the EBMT registry indicate on a total of 1714 patients autografted after 1995 in first remission (CR1) a leukemia free survival (LFS) at 5 years of 46 ± 2% highly reproducible and indeed identical when comparing Eastern Europe country data to other European countries. Several randomized studies, although not all, comparing allogeneic transplants in patients with HLA matched siblings to autologous bone marrow transplantation and to conventional chemotherapy, have shown the superiority of the allogeneic transplant approach (when feasible) to the other approaches, but also the superiority of autografting over conventional chemotherapy. None has ever shown the superiority of conventional chemotherapy. However, when reanalyzed by cytogenetics the US intergroup and the British MRC studies have shown in fact the superiority of allogeneic transplants in poor risk patients, and the superiority of ASCT in good risk patients.

The EBMT has recently investigated the outcome of patients with AML who could be defined as good risk either by clinical criteria (age <35 years and complete remission achieved within 40 days) or by cytogenetics (core binding factor mutations, inv 16 or t(8;21)) submitted to ASCT:

1) 458 adult patients with clinical good risk criteria autografted in CR1 were compared to 2218 patients classified as non good risk: the LFS was 56+/−2% versus 38+/−1%. The relapse rate was 40+/−2% versus 55+/−1%.

2) 383 patients in the EBMT registry, with inv 16 or t(8;21) were transplanted after 1990, 158 autografted and 140 allografted in CR1. Allografted recipients were younger (34 years versus 41, p< 10−4) and received their transplant earlier (Interval from diagnosis to transplant: 137 versus 161 days, p< 10−4). In addition the allograft procedure used more marrow as a stem cell source (69% vs 28%, p< 10−4) and total body irradiation (60% versus 26%, p< 10−4) rather than myeloablative chemotherapy in the conditioning. In CR2, 32 patients were autografted and 52 allografted.

Interestingly in CR1 LFS was similar following both transplant procedures (allografts: 61 ± 5%, autografts 56 ± 5% at 10 years). In contrast in CR2 allografting resulted in superior outcome (LFS : 58 ± 7% versus 30 ± 11%).

The non relapse mortality following the autograft procedure was only 5 ± 4% in CR1, but 17 ± 11% in CR2.

For patients in CR1, the median age of the population was 37 years. In those below 37 years, the LFS following allo and autografting were respectively 73 ± 6% and 58 ± 7%. In those above 37 years the results were 52 ± 7% and 60 ± 7% suggesting that autografting may be safer in older patients with core binding factor mutations.
Recent studies from the Pethema group and from UCSF confirm these findings. In the Pethema LMA 99 protocol, the LFS following ASCT in adult AML was 53% at 4 years, but in fact around 60% in patients with good and intermediate groups versus 23% only in patients of the poor risk category. In UCSF 9302 protocol, the DFS for all patients was 52% at 12 years, but in fact 68% in patients with favorable cytogenetics, 48% in patients of the intermediate risk category and 10% only in the poor risk category.

These data highlight the fact that ASCT in AML is most likely to benefit, as in other malignant blood diseases (lymphomas in particular) to patients with good prognostic criteria including high chemosensitivity. There is a need in this good risk patient population to launch randomized studies comparing conventional chemotherapy including high dose ARA-C to ASCT.

These results also are consistent with the recent EBMT analysis of of 625 patients with acute promyelocytic leukaemia (APL M3) transplanted with auto- or allogeneic-HSCT after 1993. Estimated 5 years-leukemia free survival for patients transplanted in CR1 was 69% for 149 patients autografted and it was 68% for 144 patients allografted. However The reasons why these patients in CR1 were transplanted remain unclear in the ATRA era.

For transplants in CR2, 5-y LFS was 47% in 195 autoHSCT and 59% in 137 alloHSCT recipients, respectively. ASCT is an important therapeutic tool in patients with M3 AML achieving molecular CR2.

An important question is whether adult patients with AML and no family matched donor should go to ASCT or to unrelated transplants. The Center for International Blood and Marrow Transplant Research has recently compared ASCT to unrelated donor allotransplants: they studied the outcomes of 668 autotransplants compared with 476 URD transplants. Proportional hazards regression adjusted for differences in prognostic variables. In multivariate analyses, transplant-related mortality (TRM) was significantly higher and relapse lower with URD transplantation. Adjusted 3-year survival probabilities were: in CR1 57 (53-61)% with autotransplants and 44 (37-51)% with URD (P = 0.002), in CR2 46 (39-53)% and 33 (28-38)% respectively (P = 0.006). Adjusted 3-year leukaemia-free survival (LFS) probabilities were: CR1 53 (48-57)% with autotransplants and 43 (36-50)% with URD (P = 0.021), CR2 39 (32-46)% and 33 (27-38)% respectively (P = 0.169).

Both autologous and URD transplantation produced prolonged LFS. High TRM offsetted the superior antileukaemia effect of URD transplantation. The conclusion was that this retrospective, observational database study showed that autotransplantation, in general, offered higher 3-year survival for AML patients in CR1 and CR2. Cytogenetics, however, were known in only two-thirds of patients and treatment bias could not be eliminated.

The recent introduction of non myeloablative transplants with a reduction of TRM has reinitiated the debate and rendered the decision tree more difficult to build.: The EBMT registry has compared retrospectively the outcome of 204 HLA-identical sibling RIC allo transplants (RIC) versus 954 auto transplants done from 1997 to 2003 in patients over 50 years of age. For RIC 87% of the non myeloablative regimens were built around fludarabine. For ASCT, the conditioning contained Total Body Irradiation (TBI) in 35% of the cases. In RIC patients the incidence of acute graft versus host disease (GVHD) score III-IV was only 9% but the cumulative incidence of chronic GVHD at 1 year was 46 ± 4 % (50% extensive). The non relapse mortality at 2 years was 20±3% following RIC versus 11±1% following ASCT. The relapse incidence was higher following ASCT in CR1 than following RIC (37±5% versus 25 ± 3, p= 0.03). The LFS in patients transplanted in CR1 were superposable at 2 years (43±2% following ASCT, 41±6% following RIC). The quality of life was likely better following ASCT in the absence of chronic GVHD. In CR2 result were better following RIC transplants: 57±9% versus 26±6% only following ASCT.

The question whether purging the graft in vitro improves the outcome is no longer addressed although several studies in the nineties have shown its efficacy. Part of the reason has been the long duration of aplasia following autografting with marrow treated in vitro with cyclophosphamide derivatives. Recent studies have focussed on purging peripheral blood hemopoietic stem cells with mafosfamide, and on expansion in vitro of grafts purged by mafosfamide: these studies have produced preliminary data showing that rapid engraftment can be obtained following the use of mafosfamide. In parallel, new agents for in vitro purging, that spare normal counterparts, such as TDZD-8 (4-Benzyl-2-methyl-1,2,4-thiadiazolidine-3,5-dione) are being studied.
References


Treatment of Adult ALL

Charles Linker
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**Initial Evaluation**

The initial evaluation of cases of suspected adult ALL should include histochemistry, flow cytometry, cytogenetics, and molecular testing for bcr-abl. Histochemical stains with peroxidase and esterase can help identify cases of myeloid leukemias or biphenotypic leukemias. Flow cytometry will help establish a diagnosis of ALL and will indicate whether the lineage is B or T-lymphocyte in nature. The small percentage of cases of mature B-cell ALL (Burkitt’s leukemia) will be identified by the lack of TdT and by the presence of surface immunoglobulin expression. There is also strong expression of CD20, not seen in most ALL. For the dominant group of precursor B-lineage ALL, the lack of CD10 expression can point in the direction of a pro-B ALL that may have the 11q23 cytogenetic abnormality. Of note is that almost all cases with Philadelphia chromosome-positive (Ph+) ALL co-express CD19 and CD10. Among the T-cell ALLs, it is important to identify the presence of CD2 expression. The primitive pre-thymic T-cell ALL cases have a poor prognosis, as do the mature post-thymic cases that co-express CD2 and CD3 but lack CD1a expression.

Standard cytogenetics are very important in the evaluation of ALL. Only a small fraction of adult cases will have true hyperdiploidy, as is seen in pediatric ALL, but these cases have an unusually favorable prognosis and should be identified. In the adult situation one is primarily concerned with identifying t(4,11) or other 11q23 abnormalities and identifying t(9,22) or the Philadelphia chromosome. A small percentage of cases have monosomy 7, and this is also a very high-risk patient group.

It is imperative that cases with B-precursor ALL be evaluated with molecular probes for bcr-abl, and this abnormality can be missed with standard cytogenetics. These patients require a different approach including tyrosine kinase inhibitors and should also have allogeneic stem cell transplant as part their initial therapy when possible. Among the bcr-abl-positive ALLs, the p190 abnormality is somewhat more frequent than p210 and also indicates a more aggressive disease.

**Induction therapy**

With modern induction chemotherapy at least 90% of younger adults (up to age 60) should enter complete remission. The backbone of induction therapy includes Daunorubicin, Vincristine, Prednisone, and Asparaginase. It is not clear that the addition of Cyclophosphamide adds to the effectiveness. Patients with Ph+ ALL should have imatinib or dasatinib added concurrent with initial chemotherapy, and this has made a major difference in both short-term and long-term prognosis. The remission rate has improved from 60% to 90% with the use of concurrent chemotherapy and the tyrosine kinase inhibitors. Patients over age 60 with ALL tolerate asparaginase poorly, and this should probably be omitted for these older adults.

Many adult oncologists feel uncomfortable with the use of L-asparaginase, and it is important to recognize how useful this drug has been in the
management of ALL. Early studies in pediatrics showed a significant single-agent response rate, and when it was added to vincristine and prednisone the complete remission rate increased substantially from 80% to 90%. Early studies in adults also showed an improvement in complete remission rate from 30% to 50% with the similar addition of asparaginase to vincristine and prednisone. Common toxicities of asparaginase include hyperglycemia which requires monitoring and management, as well coagulopathy which tends not to cause bleeding but rather can lead to an incidence of thrombosis. The more serious toxicities of Asparaginase include hepatotoxicity which can be severe and even fatal. There are also occasional severe anaphylactic reactions that can be quite dangerous. Recent studies in pediatrics have highlighted the efficacy of asparaginase and its important role in post-remission therapy and in increasing the long-term cure rates. A large EORTC study randomized patients between E. coli and Erwinia L-asparaginase. The Erwinia form was significantly less toxic with less neurotoxicity and less coagulopathy; however, there was a significant decrease in effectiveness with event-free survival decreasing from 73% to 60%. Pharmacokinetic investigation demonstrated that the Erwinia Asparaginase produced a more short-term asparagine depletion, four rather than eleven days, and this is probably the explanation for both the decreased toxicity and decreased effectiveness. The Pediatric Oncology Group in the United States performed a randomized study between pegylated Asparaginase either weekly or every other week. There was a significant improvement in the complete remission rate of patients treated in first relapse, 97% versus 82% (p= 0.003), with the weekly asparaginase. Pharmacokinetic studies in pediatrics have suggested an age dependence to the way Asparaginase is handled, with children over age 10 requiring 25% lower dose than younger children. The pharmacology of asparaginase in adults has not been well worked out, but it may be that pediatric doses cannot be strictly translated into adult therapy.

**Post-remission therapy**

Once remission is achieved, postremission therapy should be chosen based on a risk-adapted strategy. Approximately one-third of adult patients have a very favorable prognosis. These can be defined by the achievement of complete remission after one course of chemotherapy and the lack of adverse cytogenetics or molecular abnormalities. In addition these favorable patients are defined either as B-precursor patients who are both young (age less than 30 years) and with a low white blood count (WBC < 30,000). Thymic T-cell patients defined by expression of CD2 and usually having a mediastinal mass also have a very favorable prognosis. Alternatively a one-third of patients have a high-risk disease. These can be defined either by the requirement for more than one course of induction therapy to achieve remission, by the presence of adverse cytogenetics such as the Philadelphia chromosome, t(4,11), or monosomy 7, or by the presence of a white blood count greater than 100,000/uL in B-precursor patients. The remaining third of patients have a standard prognosis.

Favorable patients as defined above have an excellent outcome with at least a 70% cure rate with modern chemotherapy regimens. In my opinion these patients should not be treated with allogeneic transplantation in first remission. Poor risk patients by definition fare extremely poorly with chemotherapy, and there is no reasonable expectation of cure. These patients should be treated with allogeneic stem cell transplantation in first remission when possible.

The optimal therapy for standard risk adults with ALL (under age 60) remains to be defined. Most large trials of chemotherapy have suggested an event-free survival of approximately 35% in these patients. The results with allogeneic transplant in first remission appear superior to this and are in the range of 50%. However, it is possible that improved chemotherapy regimens could produce comparable outcomes to those seen with allogeneic transplantation. The UCSF 8707 program has produced 10-year event-free survival close to 60% in this patient group, and these results are similar to those seen with allogeneic transplantation.

In searching for ways to improve the postremission therapy of adults with ALL, several lines of investigation are possible. There has been suggestion that increasing the dose intensity of daunorubicin may improve outcomes, but this has not yet been rigorously tested in prospective trials. The importance of using asparaginase and not deleting this from the regimen in response to manageable toxicities has already been mentioned. Pediatric studies have demonstrated that pulse dexamethasone has major advantage compared to prednisone. There has been an improvement in the control of CNS disease possibly based on the
better CNS penetration of dexamethasone. Overall event-free survival has also improved in the pediatric population, but this has not been directly tested in adults. Another possibly major advantage of pulse dexamethasone over prolonged exposure to prednisone is the reduction in the incidence of late-complication avascular necrosis.

Nelarabine has recently been approved in the U.S. for the treatment of relapse patients with T-lineage disease. The safety of incorporating nelarabine into up-front therapy has been demonstrated in pediatric studies, but has not yet been tested in adults. It is possible that the addition of this agent to up-front therapy could improve the outcome for patients with T-cell disease.

One of the simplest and possibly most effective ways to improve outcomes of therapy for adults with ALL is to adhere to the principles of dose density and to avoid treatment delays. The comparison of the treatment of adolescent ALL between those treated with adult and pediatric regimens has shown a startling difference in outcomes with cure rates in the range of 35% to 40% for adults and 65% to 70% in pediatrics. Although there are many possible explanations for this difference, it is likely that the more rigorous adherence to schedule in the pediatric population plays a large role in this difference.

**Autologous transplant in ALL**

The role of autologous stem cell transplantation in the management of ALL remains to be defined. Randomized studies have not shown an advantage of autologous transplant over conventional therapy. However, it is possible and even likely that strategies for autologous transplantation can be improved so that outcomes may be improved. At UCSF we have collaborated with investigators at both Stanford and the City of Hope to develop a new strategy for autologous transplantation in ALL. This relies on intensive pre-transplant consolidation therapy with high-dose cytarabine (2000 mg/m2 bid x 4 days) and etoposide (40 mg/kg CIVI over 4 days) and a collection of peripheral blood stem cells early during the hematologic recovery from this chemotherapy. Although it was hypothesized that this would also serve as a form of in vivo purging, during early years of this protocol we added an antibody-based in-vitro purging to the regimen. One of the important components of our approach was the use of a very intensive preparative regimen combining high doses of total-body irradiation (1320cGy) with high-dose etoposide (60 mg/kg) and Cyclophosphamide (100 mg/kg). This is a regimen which is too toxic to be used in allogeneic transplantation, but it is manageable in the autologous setting. At this time we have treated a total of 30 patients, either very-high-risk patients in first remission as defined above or patients in second remission. With median follow-up of four years, five-year event-free survival is 44%. We are particularly gratified by the excellent outcome of patients with Philadelphia chromosome-positive ALL in whom a small number of patients have a 70% event-free survival.

**Conclusions**

In summary improvements in the treatment of adult ALL are needed on many fronts. It is possible that non-transplant chemotherapy regimens may be improved, and it is also possible that new strategies for autologous transplantation may define a role for this treatment plan.
The prognosis of Acute Lymphoblastic Leukaemia (ALL) in children has significantly improved with the use of modern therapeutic protocols. Currently, about 80% of children with ALL are cured with chemotherapy alone. Stem Cell Transplantation (SCT) plays an important role in patients with very high-risk (VHR) ALL in first remission or second complete remission (CR). Unfortunately, 70% of patients who might benefit from this therapy lack an HLA-matched sibling donor, and HLA polymorphism is still a major obstacle in finding a fully matched unrelated donor (UD) for 40% of the patients for whom the search for an UD is activated. That is why several institutions have recently explored alternative sources for SCT, such as unrelated Umbilical Cord Blood (UCB) or mismatched relatives (i.e. Haploidentical Transplantation: HT).

**Stem Cell Transplantation (SCT) in First Complete Remission (CR1) ALL**

Children with VHR CR1 ALL, as defined in Table 1, benefit more from related donor SCT than from chemotherapy. The gap between the two strategies increases as the risk profile of the patient worsens.

This is related to a higher relapse rate in children lacking a matched sibling donor (MSD), as compared to children with a MSD.

**Stem Cell Transplantation (SCT) in Second Complete Remission (CR2) ALL**

The I-BFM Study Group has defined Indications for allogeneic SCT in CR2 ALL, on the basis of the site and timing of the relapse (Table 2). DFS of patients given a MSD SCT following early relapse is significantly higher, as compared to chemotherapy. For those experiencing late relapse, the difference does not reach statistical significance.

**Unrelated Donor Stem Cell Transplantation (UD-SCT)**

Over the last 25 years, more than 18,000 UD-SCTs have been performed worldwide and facilitated by a network that includes more than 11 million volunteer UD donors enrolled in 89 registries. Currently, a suitable donor is located for 85% of the patients for whom a search is activated, and 70% of the donor phenotypes are found more than 4 times. The outcome of UD-SCT correlates with HLA matching: a single Class I or a single Class II mismatch is not relevant; multiple Class II mismatches are better than multiple Class I mismatches, which are better than Class I plus Class II mismatches. Presently, the outcome of children with CR2 ALL given an UD SCT is comparable to that of SCT from MSDs. This improvement is mainly due to refinements in HLA typing, GvHD prophylaxis and supportive care.

**Unrelated Umbilical Cord Blood Transplantation**

Throughout the last 7 years, 373 Transplant Centres in 43 countries have performed more than 3,000 UCBTs by means of a network that includes more than 130,000 cord blood units in 37 banks. In children, the results of UD bone marrow or UCBT are similar; however, types of complications differ, with more GvHD being observed in the UD
SCT group and more early deaths in the UCBT group. UCBT is a reasonable option when there is no HLA identical donor available.4

Haploidentical Transplantation

Few reports are available regarding the use of haploidentical transplantation for childhood ALL. This approach offers a promising treatment option for children with ALL requiring an urgent transplantation but lacking a suitable donor.3,11

Autologous Stem Cell Transplantation

ABMT is an effective treatment modality after early, isolated, extramedullary relapse, however only a few patients survive after late bone marrow relapse.12

Conclusions

A summary of the results that have been achieved by various strategies are reported in Table 3. We suggest that when a patient is found to have an indication for SCT, HLA typing of the patient, of the parents and siblings must be performed, including the study of ABC loci with median resolution and of DR and DQ loci with high resolution techniques. The ABO group must be studied as well. If a MSD or a phenotypically matched or 1-antigen mismatched relative is available, the patient should proceed to transplantation as soon as possible. Otherwise, the patient should be HLA-typed by high-resolution testing and a simultaneous search for an UD and an UCB should be started. A crucial point is the impact of the marrow UD search duration on the outcome of children with CR2: relapse during the search is the main limiting factor for the success of UD SCT in children with CR2 ALL.13 On the basis of whether an UD or an UCB is available within 3 months from search activation, one of the 2 options must be offered or, as an alternative, haploidentical SCT should be performed.

A further improvement in results of transplants from alternative donors should be achieved by an extensive and accurate HLA typing (allele level: A, B, C, DR, DQ, DP) to select the best match among the alternative sources of SC donors, by reducing the relapse incidence (monitoring minimal residual disease and chimerism, modulation of GvHD prophylaxis and treatment) and by reducing toxicity and infections through homogeneous supportive care and monitoring of viral and fungal infections.

Future goals include: improving results of transplants from alternative donors through precise HLA typing and homogeneous donor selection reducing relapse incidence, shortening GvHD-prophylaxis, and monitoring minimal residual disease and chimerism; reducing toxicity and infections through homogeneous supportive care and monitoring of viral and fungal infections.

Table 1. Indications for CR1

<table>
<thead>
<tr>
<th>Very high risk</th>
<th>MSD</th>
<th>UD</th>
<th>Haplo</th>
</tr>
</thead>
<tbody>
<tr>
<td>PPR &amp; t (9:22)</td>
<td>+</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>NR day 33</td>
<td>+</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>MRD day 77: &gt; 102</td>
<td>+</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>Pro-B-ALL</td>
<td>+</td>
<td>+</td>
<td>-</td>
</tr>
<tr>
<td>M3 day 15 (except T)</td>
<td>+</td>
<td>+</td>
<td>-</td>
</tr>
<tr>
<td>WBC &gt; 100,000 /μl (except T)</td>
<td>+</td>
<td>+</td>
<td>-</td>
</tr>
<tr>
<td>T-ALL: siblings only</td>
<td>+</td>
<td>+</td>
<td>-</td>
</tr>
<tr>
<td>Hopefully benefit</td>
<td>+</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>PGR &amp; t(4;11)</td>
<td>+</td>
<td>-</td>
<td>-</td>
</tr>
</tbody>
</table>

MSD, matched sibling donor; UD, unrelated donor; Haplo, haploidentical; BM, bone marrow; PPR, prednisone poor responders; MRD, minimal residual disease; NR, non responders; WBC, white blood cell count; PGR prednisone good responders

Table 2. Indications for CR2, > CR2

<table>
<thead>
<tr>
<th>Very High Risk</th>
<th>MSD</th>
<th>UD</th>
<th>Haplo</th>
</tr>
</thead>
<tbody>
<tr>
<td>all T-phenotypes</td>
<td>+</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>Non T: Very early BM</td>
<td>+</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>Or early</td>
<td>+</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>(&gt; CR 2)</td>
<td>+</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>High risk</td>
<td>+</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>non T: Early combined MRD &gt; 103</td>
<td>+</td>
<td>+</td>
<td>-</td>
</tr>
<tr>
<td>Late BM</td>
<td>+</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>all T (9:22)</td>
<td>+</td>
<td>+</td>
<td>-</td>
</tr>
<tr>
<td>&quot;Standard risk&quot;</td>
<td>+</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>non T: Late BM</td>
<td>+</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Late combined MRD &lt; 103</td>
<td>+</td>
<td>-</td>
<td>-</td>
</tr>
</tbody>
</table>

MSD, matched sibling donor; UD, unrelated donor; Haplo, haploidentical; BM, bone marrow; MRD, minimal residual disease

Table 3. Results of various strategies

| Allo SCT from MSD | Superior to CT in high risk patients |
| Allo SCT from UD | Comparable to Allo SCT from MSD |
| T-cell depletion | Promising treatment option |
| UCB transplantation | Comparable to UD SCT |

Allo SCT, allogeneic stem cell transplant; MSD, matched sibling donor; CT, chemotherapy; UD, unrelated donor; UCB, unrelated cord blood transplant
References


Several recent reports have indicated that not only a shift in the aetiology of infections and resistance patterns in patients with febrile neutropenia, but also important differences between regions and countries. Viridans streptococcal bacteraemias are common among cancer patients being second only to the coagulase-negative staphylococci. However, in certain centres in Europe Gram-negative bacilli have once again become the predominant infecting pathogens. The problems associated with emerging resistance have been widely documented in the literature. In some institutions methicillin-resistance among coagulase-negative staphylococci has reached very high proportions, and in others the incidence of extended-spectrum beta-lactamase producing Gram-negative bacilli has risen markedly. These shifts in antimicrobial susceptibility are important in guiding the choice of agents for febrile neutropenia. Antibiotic use and prophylaxis have both been associated with changes in susceptibility, and prescribing habits may influence emerging resistance. In this context, the choice of empirical antibiotic therapy and the use of prophylaxis should be driven by a sound understanding of local circumstances.

Initiating empirical broad-spectrum antibacterial therapy has long been the standard practice for febrile neutropenic cancer patients. However, during the last decade it has become evident that patients with febrile neutropenia do not constitute a homogenous group. The risk factors for developing infection and other major complications vary widely in different subsets of patients with cancer. Therefore, a valuable risk assessment of every febrile neutropenic patient is essential in order to define a tailored therapeutic approach. Those patients with hematological malignancies and severe and prolonged neutropenia will fall into the category of “high-risk”, while others who were treated with less intensive chemotherapies and who were expected to have a short duration (e.g. less than 7-10 days) of neutropenia and fewer complications during the course of neutropenia will be categorized in the ‘low-risk’ group. Recently published “The Multinational Association for Supportive Care in Cancer (MASCC)” risk index has been shown to be a valuable tool for identifying low-risk patients among adult febrile neutropenic cancer patient population. Patients with solid tumors who were treated with conventional chemotherapy and with minimal or no comorbidities (such as mucositis, cellulitis, anorectal infection, pneumonia) will usually be placed into the category of “low-risk”. On the other hand, more intensive chemotherapies have been increasingly used in solid tumor patients and some of them will also undergo an autologous hematopoietic stem cell transplantation (AHSCT). This approach will obviously increase the expected duration of neutropenia, the incidence of other comorbidities (e.g. mucositis), and may also affect the hemodynamic and clinical stability of the patient.

Once the patient is stratified in one of the risk groups, several options for empirical treatment exist. Nevertheless, several other factors need to be considered regarding to specific antimicrobial regimen. Among these are local epidemiological pattern of the infecting microorganisms and their
antimicrobial resistance pattern. Recent published data indicate that low-risk patients who are able to swallow can successfully be treated with oral antibiotics. The most frequent used regimen for this indication is a combination of a quinolone derivative (e.g. ciprofloxacin) and amoxicillin/clavulanate. Newer quinolones with enhanced activity against gram-positive pathogens (e.g. moxifloxacin, gatifloxacin) have been currently under evaluation for a monotherapy option. This type of therapy is applicable for both inpatient and outpatient settings. Stringent criteria need to be applied for selecting patients who will be treated in an outpatient program which also requires a strong commitment from both patient and healthcare team’s side. Another option is to admit the patient to the hospital and treat with parenteral antibiotics until defervescence, and then switch to oral therapy. This provides a viable alternative for patients receiving more intensive chemotherapy for treating cancer with or without AHSTC. Upon switch to an oral regimen the patient could be discharged if his/her clinical condition is permissive. Comparative solid data for such a practice are lacking yet in the literature, however several studies both in IALTG/EORTC and in elsewhere are being undertaken on this issue. For the initial parenteral therapy, monotherapy with various beta-lactam antibiotics has been extensively studied comparing with different beta-lactam plus aminoglycoside combinations. The data indicate that monotherapy with a broad-spectrum cephalosporin (e.g. ceftazidime, cefepime) or beta-lactam/beta-lactamase inhibitor combination (e.g. piperacillin/tazobactam) or a carbapenem (i.e. imipenem or meropenem) is as effective as a beta-lactam plus aminoglycoside combination for initial empirical regimen. Specific concerns for ceftazidime use exist since this drug has been held responsible for increased incidence of extended-spectrum beta-lactamase producing klebsiella infections in some institutions. Recently published metaanalyses caused concern about cefepime which was found to cause increased mortality in patients due to unexplained reasons. Parenteral quinolones has been less studied for this indication and the data are inconclusive. Therefore quinolones can not be recommended as the initial parenteral agent.

Glycopeptides should not be incorporated into the initial empirical regimen, until a document-ed gram-positive bacterial infection is observed. Recent data indicate that empirical addition of these agents is also unnecessary in those patients without defervescence after 60-72 hours of empirical broad-spectrum antibacterial therapy. Actually, glycopeptide use should strongly be discouraged unless the patient has a documented gram-positive bacterial infection or has strong predisposing factors to acquire such infections (e.g. clinically documented vascular catheter infection, colonization with methicillin resistant staphylococci or penicillin resistant pneumococci).

In summary, a risk-based approach in patients with febrile neutropenia could be more cost-effective. Various regimens with different antibiotics are available, but specific regimens also need to be tailored to local epidemiological factors.

References
Follicular Lymphoma (FL) Pathology

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Follicular lymphoma (FL) is the most common type of low-grade B cell lymphoma seen in western countries. It is characterized by a clinically indolent course. The cellular origin of the neoplastic cells are follicular center B lymphocytes [1]. The incidence of FL in eastern countries is low.

FL predominantly involves the lymph nodes, but spleen, bone marrow, peripheral blood, and Waldeyer’s ring involvement have also been reported. The gastrointestinal tract, soft tissue and skin are the most commonly involved extranodal sites.

Histologically, FL is composed of centrocytes and centroblasts, and usually has a follicular growth pattern. Neoplastic follicles are often ill-defined, and lack mantle zones. When the mantle zones are preserved there could be difficulty on differentiation form reactive follicular hyperplasia. Interfollicular infiltration of the neoplastic cells is a helpful diagnostic criterion for the cases having this morphology. Diffuse pattern may be seen and it is thought to be of clinical significance. In the WHO classification, FL is graded as 1, 2, 3a, and 3b according to the number of centroblasts per high-power field [1]. Histological grade correlates with prognosis in FL, with grades 1 and 2 being indolent and grade 3 being more aggressive. In grade 3 FL, the presence of a diffuse component is commonly seen and some studies have demonstrated that this finding is correlated with a worse outcome [2]. Presence of residual reactive follicles within the involved lymph nodes could reflect the stage of the disease [3].

Cases of ‘in situ localization of FL’ have been reported in the literature [4]. It appears to represent early microscopic involvement of FL within the lymph nodes. The clinical significance of these cases without other evidence of lymphoma is not known yet [4].

The tumor cells are positive for CD19, CD20, CD22, CD79a, surface Ig (IgM+/IgG-, IgG or rarely IgA), bcl-2, CD10, and bcl-6 and negative for CD5, CD43, CD23 and Cyclin D1 [1]. Immunohistochemistry is very useful for the diagnosis of FL, and several studies revealed the relation of expression of various proteins with clinical outcome. The proliferation index of the cells within the neoplastic follicles by MIB-1 (Ki-67) provides a measure of proliferative rate, and has been shown to correlate with FL grade but has limited prognostic significance in some studies. Some recent data revealed that proliferation index may have prognostic value in FL [5]. High Ki-67 staining in the reactive lymphoid follicles is useful for the differentiation of reactive follicular hyperplasia and FL. The proliferation index of the neoplastic follicles in low grade FL (grades 1 and 2) is lower than in reactive follicular hyperplasia and grade 3 FL. But in a recent study of Wang et al., high proliferation index in low grade FL was determined in nearly 20% of their cases. The clinical behavior of these low grade FL cases showing high proliferation index was correlated with inferior disease-specific survival but higher five-year disease-free rate similar to grade 3 FL [6].

Although most patients with FL overexpress Bcl-2 protein, higher levels of expression have been correlated with worse outcome. In contrast, higher levels
of expression of germinal center markers including CD-10, Bcl-6 and PU.1 have been correlated with a favorable outcome. The presence of more than 15 CD68+ macrophages per high power field has also been shown to predict for a poor outcome.

The genetic hallmark of FL, t(14;18)(q32;q21), which juxtaposes the bcl-2 gene with the IgH gene, is seen in 80-90% of FLs. It is not associated with the prognosis. Bcl-2 protein is expressed in the majority of the cases, and its expression reduces as histological grade increases. Although FL is rarely seen in pediatric patients, it should be noted that bcl-2 expression in pediatric FL is relatively infrequent in contrast to its adult counterpart.

Primary cutaneous follicle center cell lymphoma is a variant of FL and is often bcl-2-negative as well.

A number of cytogenetic abnormalities have been described in FL, including p53 mutations, loss of p16, upregulated MYC expression resulting from translocation or other mechanisms, gains of chromosome arms 7p or 7q, Xp, 12q and 18q, as well as losses on 6q and possibly mutations of bcl-2 and/or bcl-6 genes. The presence of additional genomic aberrations, in particular 17p and 6q deletions, is more frequent in grade 2 and 3 FL patients and correlated with shorter survival and a higher rate of transformation into diffuse large B cell lymphoma.

Approximately 25-35% of FL cases transform into diffuse large B cell lymphoma as well as Burkitt’s lymphoma, precursor B lymphoblastic lymphoma and classical type of Hodgkin’s lymphoma.

### References


Follicular lymphoma (FL) is a heterogeneous disease, which is still considered as incurable. New prognostic factors as the FLIPI (Follicular Lymphoma International Prognostic Index) and histological grading are of help in deciding type of therapeutic option. Also immune cells in the micro environment of the tumour have shown to be of clinical importance.

Improvement in therapy has been seen with the monoclonal anti-CD20 antibody rituximab, first used as single agent therapy in relapsed follicular lymphoma (the pivotal trial). Recently the combination of this antibody with chemotherapy (CVP, CHOP or fludarabine combinations) has been shown to extend both progression-free (PFS) and overall survival (OS), when used as first-and or second line therapy.

To improve the efficacy of rituximab and to delay the need of chemotherapy, different schedules of drug administration, as extended dosing, combination with interferon and maintenance, has been used.

R-CHOP (rituximab 375 mg/m² at day 1 of each cycle of CHOP) was used in a phase III clinical trial for patients with relapsed follicular lymphoma, the EORTC 20981 Intergroup Study. Patients with a CR or PR after six cycles of therapy underwent a second randomisation to no further treatment (observation) or maintenance treatment with rituximab (375 mg/m² once every 3 months) until relapse or for a maximum of 2 years. Both treatment arms yielded similar PR rates but a highly significant higher CR rate with the combination. In patients randomised to maintenance an advantage was observed both in PFS and OS when compared with the observation arm.

The benefit of rituximab maintenance for patients with first-line immunochemotherapy will be determined in ongoing randomised studies.

The best schedule and length of therapy as well as longterm effects of rituximab treatment needs to be evaluated. New monoclonal antibodies are under development, but it is still not defined which subgroups of patients will benefit most of treatment with antibodies. Also other new drugs acting on the microenvironment of the lymphoma might show therapeutic activity.
Gene expression profiling has defined major subtypes of aggressive non-Hodgkin’s lymphomas (B-NHL) in recent years, in particular among diffuse large B-cell lymphomas (DLBCL). Specifically, a germinal center B-cell type (GCB DLBCL), an activated B-cell type (ABC DLBCL) and primary mediastinal B-cell lymphoma (PMBL) can be distinguished based on fundamental differences in underlying gene expression profiles. More recently, the Lymphoma and Leukemia Molecular Profiling Project (LLMPP) (Dave et al., NEJM 2006) as well as a major study conducted in Germany (Hummel et al., NEJM 2006) defined the highly aggressive entity of Burkitt lymphoma on a molecular level. Besides the molecular classification of B-NHL, gene expression signatures can also be used to predict clinical outcome. For example, the gene expression-based measurement of proliferation (proliferation signature) provides a powerful predictor of outcome in mantle cell lymphoma (MCL) that is superior to the immunohistochemical assessment of the Ki-67 index (Rosenwald et al., Cancer Cell 2003). How can these findings be translated into daily clinical practice?

Several attempts have been made to use a small set of immunohistochemical markers as surrogates for gene expression signatures. An example is the Hans-classifier (Hans et al., Blood 2004) that uses the immunohistochemical markers CD10, BCL6 and MUM1 to classify DLBCL into the GCB- and non-GCB subtypes. While this approach appears to be easily applicable in the daily routine, there are also limitations, and a considerable variation in the staining process (between different institutions) as well as in the scoring process (between different pathologists) have to be overcome. In particular, the germinal center-associated marker BCL6 appears to suffer from remarkable inter-laboratory variation (de Jong et al., JCO 2007).

Alternatively, quantitative RT-PCR (TaqMan) can be performed in routinely obtained formalin fixed and paraffin embedded tumor tissues. In mantle cell lymphoma (MCL), the quantitative mRNA expression measurement of 5 genes may be able to substitute for the proliferation signature determined by gene expression profiling (Hartmann et al., unpublished). Finally, the microarray platform itself may be used for diagnostic and prognostic purposes in clinical settings. Towards this goal, efforts are under way to achieve reliable and robust gene expression measurements across different laboratories. In one of these efforts, the Lymphoma and Leukemia Molecular Profiling Project (LLMPP) collaborates with Roche Diagnostics in the development of a diagnostic microarray. As a first step, a proficiency test between the 8 participating institutions was undertaken which showed highly consistent gene expression results of the same tumor tissues analyzed in a decentralized fashion.
The aggressive non-Hodgkin’s lymphomas can be cured in more than half of the cases. For patients with localized aggressive non-Hodgkin’s lymphoma, heterogeneity in patients selection prevent us from defining a new standard of care. On the contrary, for patients with advanced stages of diffuse large B cell lymphoma a new standard of therapy now exist particularly for elderly and low risk young patients.

In this article I will separate the patients into 3 different groups: I) early stage NHL, II) advanced stage elderly and low risk young patients and III) young high risk patients.

I. Early (Limited) Stage Aggressive Lymphoma

There is extreme heterogeneity within the group of patients described as having early stage lymphoma, making comparisons of outcome difficult.

SWOG compared 3 cycles of CHOP with RT to 8 cycles of CHOP in patients with localized aggressive NHL and found that CHOP(3) plus RT was superior to CHOP (8) through the first five years of follow-up (1) . Localized disease was defined as stage I and non-bulky stage II disease. Patients with bulky-stage II disease are known to have a prognosis similar to patients with advanced disease and should be excluded from limited-stage disease (2). Patients with bulky-stage II disease accrued to SWOG studies had a 5-year survival of 49%. Patients with stage III-IV disease had a 5-year survival of 46%. Therefore bulky stage II is considered as “advanced” disease. If patients with bulky-stage II disease are included in trials and treated with a short-course of chemotherapy plus RT they have an inferior outcome compared to similar patients treated with aggressive chemotherapy designed for advanced disease.

Reyes and colleagues from GELA (3) have reported the results of a randomized trial comparing CHOP (3) plus RT to an aggressive combination chemotherapy called ACVBP (LNH-93-1). ACVBP chemotherapy was originally designed for advanced disease. In the subgroup of bulky-stage II disease, treatment with ACVBP was superior to CHOP(3) plus RT with 5-year survival estimates of 82% and 50%, respectively (p=0.03)

By excluding patients with bulky-stage II disease one might presume that the remaining patients with limited disease comprise a homogeneous group with regard to prognosis and choice of optimal therapy. But 10 year survival can vary from 90% to 10% within subgroup of limited stage patients (4). Predicting such variable outcome easily accomplished using stage modified IPI. Most clinicians uses IPI and it includes five risk factors including age, stage, serum LDH, performance status(PS), and the number of extranodal sites of disease(5). Age, stage, serum LDH, and PS each predict significant outcome differences for patients with limited diseases. Patients with no adverse risk factors have a very good prognosis with 10 year survival estimates exceed 90% treated (Table 1 and 2). These very good results can be achieved regardless of the treatment strategy chosen; CHOP (3) +RT, CHOP (8), or ACVBP (1,4,6). This category is called ‘very limited disease’.
Another GELA study was published by Fillet et al. comparing 4 cycles of CHOP to CHOP plus 40 Gy involved-field RT in patients older than 60 years with no adverse risk factors according to the age-adjusted IPI (7). There was no advantage of the radiation and, a possible disadvantage in patients older than 70 years of age. Recent ECOG study suggested benefit of adjuvant radiation after 8 cycles of CHOP (8). But the main difference in these two trials was bulky disease rates. Although SWOG included only 6 (1.5%) bulky (stage I) patients, the rate of bulky disease in ECOG was 31%. So one third of the patients actually should be accepted as advanced disease in ECOG trial.

The MINT trial included some patients with early stage disease and compared a CHOP-like chemotherapy regimen to the same regimen with the addition of rituximab in young, good prognosis patients (9). For the most favorable patients (those without bulky disease) the results with a complete course of chemotherapy plus rituximab alone without radiation led to survival in excess of 90%. The Southwest Oncology Group in the United States reported a pilot study of an abbreviated course of CHOP plus rituximab followed by radiation showing progression free and overall survival in excess of 90% (10).

II. Treatment Of Advanced Stage Elderly And Low Risk Young Patients

After initial staging bulky stage II, stage III or stage IV aggressive lymphoma is documented in approximately 75% of all DLBCL patients. Therefore chemotherapy is main treatment modality for these patients.

The study that defined chemotherapy as standard was an intergroup trial conducted by SWOG and ECOG. In this study previously untreated patients with stage II bulky, III, and IV disease with intermediate –or high-grade histology were randomized to one of four treatment arms: CHOP, m-BACOD, ProMACE-CytaBOM or MACOP-B. There was no difference in any treatment arm. These results along with the fact that CHOP was cheaper and easier to administer than the other regimens, established CHOP as the standard therapy throughout the world. However, with a projected disease-free survival rate of 36%, it is obvious that it is far from an ideal therapy, and there is need for better treatment approaches (11).

The International non-Hodgkin Lymphoma Prognostic Factors Index (IPI) utilized pretreatment prognostic factors in a sample of over 5000 patients to develop a predictive model of outcome for aggressive non-Hodgkin’s Lymphoma (12). Five pretreatment characteristics were found to be independent predictors of death: age (< 60 vs. >60), tumor stage II or II vs. III or IV, the number of involved extranodal sites (< 1 vs. >1), patient ECOG performance status (0,1;ambulatory vs. >2 ;non-ambulatory), and serum LDH level(normal or elevated). Each of the individual factors had comparable relative risks. The resulting model identified 4 risk groups with the following associated 5-year survival rates: low risk(0-1), 73%, low-intermediate risk (2 risk factors), 51%; high-intermediate risk (3 risk factors), 43%, and high risk (4-5 risk factors), 26%. However, the improvement in treatment response associated with the addition of the antibody rituximab to treatment regimens seems to have altered survival of prognostic groups using the International Prognostic Index (13).

<table>
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<tr>
<th>Table 1. International Prognostic Index (12)</th>
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<tr>
<td>Full Index</td>
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<tr>
<td>Age adjust</td>
</tr>
<tr>
<td>Prognostic Factors</td>
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<tr>
<td>Age &gt; 60 years</td>
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<tr>
<td>Performance status &gt; 2</td>
</tr>
<tr>
<td>LDH &gt; 1 x normal</td>
</tr>
<tr>
<td>Extranasal sites &gt; 2</td>
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<tr>
<td>Stage III or IV</td>
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<td>Risk Category Factors</td>
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<tr>
<td>• Low</td>
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<td>Risk Category Factors</td>
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<td>• Low</td>
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<tr>
<th>Table 2. Outcome for patients with diffuse aggressive lymphoma after anthracycline chemotherapy International Index (12)</th>
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<tr>
<td>Risk Category No Risk Factors</td>
</tr>
<tr>
<td>% Cases CR rate PFS of CRs 5 yr Survival 5 yr</td>
</tr>
<tr>
<td>Low 0.1 35% 87% 70% 72%</td>
</tr>
<tr>
<td>Low-intermediate 2 27% 67% 50% 50%</td>
</tr>
<tr>
<td>High-intermediate 2 22% 55% 48% 43%</td>
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<tr>
<td>High 4.5 18% 44% 40% 26%</td>
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<th>Table 3. Results with a revised IPI when R-CHOP is given for diffuse large B-cell lymphoma (13)</th>
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<tr>
<td>Group No Factors % Patients % 4-Year Overall Survival</td>
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<tr>
<td>Standard IPI 0.1 28 86</td>
</tr>
<tr>
<td>2 27 81</td>
</tr>
<tr>
<td>3 21 54</td>
</tr>
<tr>
<td>4.5 24 58</td>
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<tr>
<td>Revised IPI 0.1 10 92</td>
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<tr>
<td>1.2 45 82</td>
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<td>3,4 45 58</td>
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Currently molecular profiling project has used complementary DNA (cDNA) microarray techniques to demonstrate two distinct subpopulations.
of DLBCL with different prognosis and different genetics. Patients with a germinal center B (GCB) cell like signature have a more favorable course than those with an activated B-cell (ABC) signature (14,15). At the present time gene expression profiling is not part of routine clinical practice (37). This may be due to technical difficulties in performing the c-DNA arrays. GCB versus ABC typing seems to be able to be reproduced by tissue microarray (16).

After the disappointing results with the third generation chemotherapy regimens, several new treatment approaches have been developed. GELA developed the ACVBP regimen (which involves very intensive chemotherapy for four courses followed by an intensive consolidation) which was shown to be superior to CHOP in subgroup of patients (17).

German high-grade lymphoma study group found that addition of etoposide to CHOP improved results in young patients (18), while CHOP administered at 14 day instead of 21 day intervals seemed to improve the results in elderly patients (19).

An infusional chemotherapy regimen developed at the National Cancer Institute referred to as DA-EPOCH had very encouraging results (20).

The randomized study that changed practice throughout the world was performed by the GELA and compared CHOP versus Rituximab-CHOP (R-CHOP) in elderly patients (21,22). The GELA group randomized 399 previously untreated patients with DLBCL, 60 to 80 years old, to receive either 8 cycles of CHOP every 3 weeks or 8 cycles of CHOP plus rituximab given on day 1 of each cycle. Complete response rates of 76% and 63% (p=0.005) and 5-year overall survival rates of 58 % and 45 % (p< 0.007) were achieved by R-CHOP and CHOP, respectively. In patients with a low-risk age adjusted IPI (0 or 1 adverse prognostic factors), 5-year event-free survival (EFS) was 63% and 34% for the R-COP and CHOP arms, respectively (p=0.0008). For those with high-risk disease (2, 3 adverse factors), 5 year EFS were 41% and 27% for the R-CHOP and CHOP arms, respectively (p=0.004). These results suggest that addition of rituximab therapy to standard CHOP may lead to significant prolongation of event-free and overall survival in elderly patients with both high-risk and low risk disease with no increase in toxicity. In an early analysis, R-CHOP appeared to be more effective than CHOP in bcl-2 positive, but not in bcl-2 negative patients, suggesting that the benefit of addition of rituximab might overcome bcl-2 associated chemotherapy resistance (23,24).

In the United States a larger (N=632) intergroup study randomized elderly patients to receive initial therapy with either CHOP or R-CHOP. The rituximab was given on a different schedule than the GELA study. Responding patients then were randomized to receive either rituximab maintenance therapy (4 doses q 6 months x 2 years) or no maintenance. This study confirmed the GELA results with a significant advantage for receiving rituximab either in during induction or maintenance, but no advantage to getting both (25).

According to the IPI young good prognosis patients comprise the low and low-intermediate risk group (0 and 1 risk factor according to the age-adjusted IPI) and poor prognosis patients the high-intermediate and high risk group (2 risk factors).

Until recently there was no Phase III trial looking for the value of rituximab in younger patients with DLBCL. An International study called MinT Trial compared chemotherapy CHOP with or without rituximab in patients younger than age 60. Eligibility criteria included DLBCL, 18-60 years, IPI 0 or 1, stages II-IV or stage I with bulk. Patients received 6 cycles of any of several CHOP like regimens followed by radiation therapy to bulky disease (9). The MinT trial demonstrated significant advantage in response, failure-free survival, and overall survival with the addition of rituximab.

The German High Grade Lymphoma Study Group (DSHNHL) studied the utility of six versus eight cycles of R-CHOP at 14 day intervals with or without rituximab in elderly patients with DLBCL. RICOVER 60 trial of DSHNHL demonstrated the importance of rituximab and in combination with rituximab 6 cycles of R-CHOP-14 are as good as 8 cycles.(26)

Investigators from the Cancer Institute from the British Columbia did a population based study of the impact of adding rituximab to CHOP. After approval of rituximab in British Columbia survival of DLBCL went up of about 20%. (27)

### III. Treatment Of Young, High Risk Patients

According to the IPI, young poor prognosis patients comprise the high-intermediate and high risk group (aa IPI 2,3). For young poor prognosis patients, the five-year survival is around 50%,
and progress has not been demonstrated in these patients.

The role of high-dose therapy (HDT) followed by autologous stem cell transplantation (SCT) for primary treatment of aggressive non-Hodgkin lymphoma is still uncertain. Whereas some studies demonstrated superiority of HDT over conventional treatment (28, 29,30), others failed to show significant differences (31,32,33) or reported inferior results (34). Besides differences in patient characteristics, the type HDT and its timing may have important implications for outcome. Recently published meta-analysis showed that there was no evidence that HDT improved OS and EFS in good risk NHL patients. The evidence for poor risk patients is inconclusive and high quality studies in poor risk patients are warranted (35).

Recently, German High Grade NHL study group published the result of Mega CHOEP chemotherapy. The Mega CHOEP protocol consisted of 4 courses of Cyclophosphamide, doxorubicin, vincristine, etoposide and prednisolone. MegaCHOEP was applied as high dose chemotherapy including autologous stem cell transplantation after courses 2, 3, 4. The patients aged 18-60 years with primary diagnosis of aggressive NHL and LDH levels above normal were included in the study. 70% of the patients achieved CR or CRu after a median follow-up of 55 months. OS was 75% and 67.2% after 2 and 5 years, respectively. Treatment related mortality was 4.5%. Risk of secondary MDS/AML was low with only one patient having MDS during follow up.

Comparison of the results of Mega CHOEP with R-CHOP chemotherapy is difficult. The only study published with R-CHOP chemotherapy was that of British Columbia’s and 4 year OS was 55% for this high risk patient group. But we have to keep in mind that the median age of this group was older (median age 61) than the patients included in the Mega CHOEP study (median age 43).

References


DLBCL-L Relapse, Resistant Cases and Transplantation

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The University of Chicago, Chicago, USA

Treatment of Relapsed and Refractory DLBCL

More than half of patients with aggressive lymphoma, initially entering remission with combination chemotherapy will relapse. The standard treatment approach for such patients is to deliver salvage chemotherapy followed by consolidative autologous stem cell transplantation in patients demonstrating chemosensitivity. Patients with chemorefractory disease and patients relapsing following an autologous stem cell transplant have an overall poor prognosis and should be considered for allogeneic stem cell transplantation or for clinical trials with investigational agents.

The optimal salvage regimen is not known, and there are no phase III prospective randomized trials comparing various combinations. Most of the data is from phase II trials, and the choice of treatment is often influenced by both patient features and physician preferences. Some commonly used regimens include DHAP (dexamethasone, cytarabine, cisplatin), ESHAP (etoposide, methylprednisolone, cytarabine, cisplatin), ICE (ifosfamide, carboplatin, etoposide), IE (ifosfamide, etoposide), MINE (mesna, ifosfamide, mitoxantrone, etoposide) and EPOCH (etoposide, prednisone, vincristine, cyclophosphamide, doxorubicin), among others. The DHAP regimen is one of the first salvage regimens to be designed. In the Parma trial, patients with relapsed lymphomas receiving DHAP had an overall response rate of 58%, but the 5 year event-free survival and overall survival of patients not subsequently transplanted were only 12% and 32%, respectively.

Ifosfamide-based regimens are gaining in popularity, partly due to the ability to escalate the ifosfamide, and also because they are excellent stem cell mobilizing regimens. Overall response rates are over 60%, although the complete response rate is only 24%. The major advantage to improving salvage regimens is to demonstrate chemosensitivity, since this is arguably the most crucial characteristic determining outcome following autologous stem cell transplantation in aggressive lymphomas. Of the ifosfamide-based salvage regimens for aggressive lymphomas, extensive data has been published on the ICE (ifosfamide, carboplatin, etoposide) regimen developed at the Memorial Sloan Kettering Cancer Center (MSKCC). In an initial publication, investigators at MSKCC treated 163 consecutive transplant-eligible patients with relapsed or refractory aggressive NHL with 3 cycles of the ICE regimen. The overall response rate was 66%, allowing 89% of patients to proceed to a planned autologous stem cell transplant. There was minimal non-hematologic toxicity, although a third of patients had greater than grade 3 thrombocytopenia. All patients received growth factor support during each cycle of treatment. There are several other high dose ifosfamide-based regimens that are in widespread use, and all appear to be effective at stem cell mobilization. However, despite high activity, none of these regimens are curative unless followed by a consolidative transplant procedure.

The addition of rituximab to salvage regimens appears to substantially improve the response rate. For example, Kewalramani and colleagues
show that the overall response rate and complete response rate increases to 81% and 55%, respectively, when adding rituximab to the ICE regimen. Although not specifically demonstrated for large cell lymphoma, rituximab also serves as an "in vivo purge" during stem cell collection [79,80] and is likely to be an important component of most pre-transplant salvage regimens for CD20-positive malignancies.

**New Investigational Drugs:**

There are a multitude of promising investigational agents being pursued for the treatment of lymphomas. These include proteosome inhibitors (bortezomib or Velcade®), anti-Bcl-2 agents (oblimersen sodium or Genasense®), anti-angiogenic agents, liposomal formulations of standard chemotherapeutic agents (liposomal vincristine, liposomal doxorubicin), newer monoclonal antibodies (epratuzumab), and radiolabelled monoclonal antibodies (ibritumomab tiuxetan or Zevalin®, tositumomab or Bexxar®). Phase II and III studies are ongoing, and several of the most active agents in preliminary studies are being incorporated into front-line regimens.

We have had a particular interest in the development of ixabepilone and of temsirolimus in the management of recurrent or refractory large cell lymphoma, both drugs have excellent efficacy in large cell lymphoma and have induced remission in otherwise refractory patients. Both have a favorable toxicity profile.

**Transplantation in the management of recurrent lymphoma**

Patients receiving autologous stem cell transplants for chemotherapy-sensitive relapsed non-Hodgkin lymphoma have significantly superior survival compared to those receiving conventional chemotherapy. For example, in one large trial 5-year survival was 46% in the transplant group vs. 12% in the group receiving chemotherapy. (The very rare patient with an International Prognostic Index (IPI) score of 0 at the time of relapse appears to do equally well with chemotherapy or transplant.) Most now accept autologous SCT as the best therapy for relapsed, chemotherapy-sensitive lymphoma. The introduction of rituximab has improved the prognosis of aggressive B-cell lymphoma and rituximab may also have a role in the peri-transplant management of patients with aggressive lymphoma. When given before transplant it may have a purging effect. It is also given post-transplant in an effort to reduce recurrence. Interestingly, patients given rituximab before or after transplant are prone to severe but transient neutropenia, the mechanism of which is poorly understood.

Allogeneic transplantation has not been as extensively utilized for aggressive lymphoma. Although relapse rates are lower than after autologous transplantation, the risk for TRM is greater after allogeneic SCT. The role of a GVL effect continues to be investigated but the aggressive growth of the tumor may not allow for the full benefit of immunotherapy. Allogeneic transplantation may be preferable in some patient subsets such as those who fail to mobilize, have marrow involvement or have failed a previous autologous transplantation. Allogeneic SCT may also be preferable for patients with peripheral T-cell lymphoma, who have high recurrence rates after autologous transplantation.

Mantle cell lymphoma is a relentless illness with high recurrence rates after CHOP-rituximab therapy. Many centers now recommend consolidation with autologous transplantation in first remission with extremely promising results. For patients with recurrent mantle cell lymphoma, allogeneic transplantation is preferred.

**References**

Hodgkin’s Diseases First Line Treatment

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Introduction
Hodgkin Lymphoma (HL) is one of the best curable cancers of adulthood. In localized stages of the disease more than 95% of patients can be cured with modern treatment strategies. In advanced stages, comprising stages IIB with large mediastinal tumors and all stages III and IV according to the Ann Arbor classification, more than 85% of patients experience long term tumor free survival and in the current cohorts of HL survivors after 15 years more individuals have died due to adverse sequelae of treatment than of Hodgkin related causes of death. Therefore current intentions for improvement aim for preserving the high cure rates while reducing the acute and long term toxicities. The definition of advanced HL varies considerably between different study groups and makes inter-study comparisons difficult.

BEACOPP in advanced stage HL
Development and results of the three BEACOPP regimens: baseline, escalated and BEACOPP –14.

HL is a very chemo-radio-sensitive disease, the pathognomonic Reed- Sternberg cell is extremely fragile upon in vitro manipulation, the primary tumor lesion consists of less than 0.1-1.0% tumor cells, the rest is reactive tissue. Hence, biologically this tumor should be responsive to a cytotoxic regimen using an optimal drug combination in an optimal time schedule and drug dosage to yield the highest tumor cell kill.
statement, it becomes obvious that the outcome of patients with advanced HL treated in most North-American or European multicenter studies with the common conventional strategies like ABVD, MOPP/ABVD or MOPP/ABV-hybrid yield suboptimal outcome results: for ABVD at 5 years FFS was 63% and OS was 82% (2).

To improve the outcome of patients with advanced stage HL one either should use new, effective drugs, however, these are not available,- or apply the existing drugs (-combinations) in a more suitable way to increase drug- intensity and drug- density.

The GHSG, in a model design on the basis of data from previous trials with the COPP/ABVD regimen, hypothesized that shortening the treatment interval would lead to a modest benefit. In 1992 the GHSG developed the BEACOPP regimen that is administered every 21 days on the basis of the COPP/ABVD alternating regime, which is administered every 28 days.. Baseline- BEACOPP is based on the same drug composition as the standard COPP/ABVD, but excludes vinblastine and dacarbazine, and adds etoposide. The dose for the baseline schedule is equivalent to COPP/ABVD.

The model further suggested that an escalation of dosage by 30% might intensify the cell kill by 10-15% (3).

In the resulting escalated BEACOPP- regimen the doses of cyclophosphamide, adriamycin, and etoposide are increased with G-CSF support.

After getting encouraging results in pilot studies (4, 5), the three-armed HD9 trial compared the escalated and the baseline BEACOPP regimen with the standard COPP/ABVD for the treatment of advanced-stage HL patients (6).

After a median observation time of 7 years, the superiority of the dose-escalated BEACOPP regimen was clearly demonstrated in terms of FFTF (BEACOPP esc.: 85%, BEACOPP baseline: 75%, COPP/ABVD: 67%) and OS (BEACOPP esc.: 90%, BEACOPP baseline: 84%, COPP/ABVD: 79%) (1). There was a higher number of secondary AML/MDS in the escalated BEACOPP arm (n=11) compared with the C/ABVD arm (n= 1), the death rate due to treatment induced toxicity was lower in the esc.BEACOPP arm (1,6% vs 1,8%). Death due to progressive HL was 1,7% for esc BEACOPP and 8,7% for C/ABVD.

Decreasing toxicity by reducing the number of escalated BEACOPP cycles and using the baseline BEACOPP-14 regimen: preliminary results of the completed HD12- and ongoing HD15- studies

Taking in account the high chemosensitivity of the fragile Reed- Sterberg cells and a very pronounced genetic lability of the tumor cells leading to early secondary resistance, it seems of outmost importance to get a maximal and rapid cell kill at the onset of therapy, possibly in the first two months after commencment treatment. Recent studies, using the PET as a predictor of response, show that an early CR is the most favorable prognostic indicator.

In North American and European multicenter studies using the conventional ABVD, MOPP/ABV etc., this aim was hardly reached when 10-15% of the patients had refractory tumors and progressed and 30-35% relapsed within 5 years, resulting in an OS rate of 70-80% at 5-10 years. The data with the escalated BEACOPP regime, however, after 7 years mot for FFTF are 85% and for OS 90%, while 99% of the AML/MDS occurred after 1-5 years, none was observed in the last two years.

Modern therapeutic strategies aim at both: reducing therapy-induced late toxicities while maintaining effective tumour control.

Therefore, in the subsequent HD12 trial of the GHSG, chemotherapy was de-escalated by comparing eight cycles of escalated BEACOPP with four escalated and four baseline cycles, with or without consolidating radiation to initial bulky and residual disease. The results are very promising and absolutely in line with the HD9 results:

After a median follow-up of 2 years, FFTF and OS for the whole cohort were 88% and 94%, respectively. For the group getting 4 escalating BEACOPP+ 4 baseline BEACOPP, FFTF was 88% and OS 94% respectively; For the patient cohort getting 8 esc BEACOPP, FFTF was 90% and OS: 96%.

In this HD-12 study less than 35% of the total cohort of patients received consolidative involved field radiation. The rate of secondary AML/MDS was at the same time point of observation only half of that in the HD-9 study.

The currently ongoing HD15 trial for the treatment of advanced-stage HL compares 8 courses of escalated BEACOPP with 6 courses of escalated BEACOPP or 8 courses of the baseline BEACOPP-14 to further reduce toxicity. As an example for
a time dense/intensified chemotherapy, the BEACOPP-14 schedule, representing a BEACOPP-baseline variant given in 14 day intervals, was developed.

Consolidative radiation was given to 65% of patients. This regimen was tested in a multicenter pilot study with the final analysis showing an estimated FFTF of 90%, an OS of 95% at 5 years median observation time. The acute hematotoxicity was significantly reduced and ranged between that of escalated and baseline BEACOPP-21 (7). In this pilot study with 94 patients after 5 years most there was no AML/MDS, no solid tumor, only 1 NHL as secondary neoplasia. There were two toxic deaths during treatment.

**Conclusion and future aims**

**The question is relevant**

Is it justified to call ABVD the goldstandard treatment strategy for advanced stage HL patients?

The answer might be yes, if one takes the recent data of North American single center studies (Vancouver) in account, where 25-30% of patients with stages I-II with bulky tumors and B-symptoms are subsummed into the group of advanced stage HL patients. These patients, however, are treated in the GHSG in the early unfavorable (intermediate) cohort with just 4 cycles of ABVD+ 20 Gy IF-RT, resulting in a 5 year FFS of 90% and an OS rate of 97%.

The robust data from the North American multicenter studies (CALGB, ECOG, Intergroup) including only stages III-IV and possibly the IIB patients with large mediastinal masses in the advanced stage group showed rather disappointing results with a 10 year FFS of 50% (8).

The answer might be no, if one accepts that at a median observation time of 7 years the GHSG data with escalated BEACOPP (HD9- study) (6, 1) showed that 11 more young patients (median age 28 years) out of a group of 100 escalated BEACOPP treated patients are alive and possibly cured, compared with the group of C/ABVD treated patients, where 21 patients had already died at this time point.

**The pivotal question is**

How much long term toxicity do we expect for escalated BEACOPP treated patients after 10, 15 and 20 years follow up? There is certainly no answer to this question at the moment. The 10 year data of the HD9 study, however, look promising and inspite of a higher number of initially occurring AML/MDS there is a significant superiority for the escalated BEACOPP regimen compared to C/ABVD and 11 more young patients survive with this admittedly very toxic principle than with the ABVD regime in a multi center study with more than 500 participating centers and many patients treated in outpatient setting, as it is the case in the GHSG.

The difference for freedom from treatment failure (FFTF) at 10 years is even higher in the two arms: 64% for C/ABVD and 82% for BEACOPP escalated. That means 18% of patients treated with BEACOPP escalated do not need a salvage treatment conferring a even higher risk of AML/MDS than BEACOPP escalated when high dose chemotherapy with autologous stem cell support is given as salvage.

To further reduce the danger of late complications, like secondary tumors, cardiac and pulmonary late toxicities, the GHSG aims at reducing the number of escalated BEACOPP cycles from a total of 8 to 6 (HD15) and in the HD12 trial even to 4 escalated BEACOPP + 4 baseline BEACOPP cycles. Parallel to this de-escalation of BEACOPP the amount of radiotherapy is reduced from 65% of patients receiving consolidating radiation in the HD9 trial to less than 5% in the HD15 trial, where involved field RT is only given to PET positive >2.5cm rest tumors. All PET negative rest tumors, irrespective of their size after 6 or 8 escalated BEACOPP- or 8 BEACOPP-14-cycles, do not get IF-RT anymore!

The pivotal question is

1. risk adapted therapy allocating the patients according to their risk of being resistant or relapsing using the International Prognostic Score (Hasenclever/Diehl) and treating patients with the lower risk IPS 0-2 with ABVD and with the higher risk IPS > 3 with BEACOPP escalated and

2. response adapted therapy using FDG-PET as a response- and prognosis predictor. That means, after 2 courses of either ABVD or BEACOPP esc and a negative PET: continue with ABVD, if PET is positive, change to BEACOPP esc.
The ultimate answer to the question whether escalated BEACOPP is superior to ABVD only can be answered in a randomized multi center, multi culture setting, testing both regimen with and without radiation and in the different strata of the risk groups of the International Prognostic Factor Project (9). A global study, lead by the EORTC is currently being undertaken in North America, Canada, Australia and in Europe (EORTC (#20012) (10).

References


Relaps and Refractory Hodgkin`s Diseases

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Introduction

Depending on stage and risk factor profile, up to 95% of patients with Hodgkin’s disease (HD) at first presentation reach complete remission (CR) after the initial standard treatment including radiotherapy, combination chemotherapy, or combined modality therapy. Patients who relapse after a first CR can achieve a second CR with salvage treatment including radiotherapy for localized relapse in previously non-irradiated areas, conventional salvage chemotherapy, or high-dose chemotherapy (HDCT) with stem cell transplantation (SCT).1

Prognostic factors in relapsed and refractory Hodgkin`s disease

It was first noted in 1979 that the length of remission to first-line chemotherapy had a marked effect on the ability of patients to respond to subsequent salvage treatment.2 In 1992 the National Cancer Institute (NCI) updated their experience with the long-term follow up of patients who relapsed after polychemotherapy.3 Derived primarily from investigations involving failures after MOPP and MOPP variants, the conclusions are thought to be relevant to other chemotherapy programs as well. On this basis, chemotherapy failures can be divided into three subgroups:

- Primary progressive Hodgkin’s disease, i.e. patients who never achieve a complete remission
- Early relapses within 12 months of CR
- Late relapses after CR lasting > 12 months

Using conventional chemotherapy for patients with primary progressive disease, virtually no patient survives more than eight years. In contrast, the projected 20-year survival for patients with early relapse or late relapse was 11% and 22%, respectively.3

Primary progressive Hodgkin`s disease

Patients with primary progressive disease, defined as progression during induction treatment or within 90 days after the end of treatment, have a particularly poor prognosis. Conventional salvage regimens have given disappointing results in the vast majority of patients: response to salvage treatment is low and the duration of response is often short. The 8-year OS ranges between 0% and 8%.3,4

The German Hodgkin’s disease Study Group (GHSG) retrospectively analysed 206 patients with PD to determine outcome after salvage therapy and identify prognostic factors.5 The five year freedom from second failure (FF2F) and OS for all patients was 17% and 26%. As reported from transplant centers, the five year FF2F and OS for patients treated with HDCT is 42% and 48%, respectively, but only 33% of all patients received HDCT. The low percentage of patients who received HDCT was due to rapidly fatal disease or life-threatening severe toxicity after salvage therapy. Other reasons not to proceed to HDCT were insufficient stem cell harvest, poor performance status and older age.
In a multivariate analysis, Karnofsky performance score at progress \( p < 0.0001 \), age \( p = 0.019 \), and attainment of a temporary remission to first-line chemotherapy \( p = 0.0003 \) were significant prognostic factors for survival. Patients with none of these risk factors had a 5-year OS of 55% compared with 0% for patients with all three of these unfavorable prognostic factors.

**Early and late relapsed Hodgkin’s disease**

The overall prognosis is bad for patients relapsing after first-line chemotherapy when treated with conventional chemotherapy. At present, HDCT followed by ASCT is the treatment of choice for patients with relapsed HD after first-line polychemotherapy. Although the results reported with HDCT in patients with late relapse have been superior to those reported in most series of conventional chemotherapy, the use of HDCT in late relapses had been an area of controversy because patients with late relapse have satisfactory second CR rates when treated with conventional chemotherapy with OS ranging from 40% to 55%. However, the HDR-1 trial of the GHSG showed improved FFTF after HDCT compared with conventional chemotherapy also in patients with late relapse.6

Many prognostic factors have been described for patients relapsing after first-line chemotherapy. These include age, sex, histology, relapse sites, stage at relapse, bulky disease, B symptoms, performance status, and extranodal relapse. The impact of these factors is difficult to assess due to confounding factors such as small number of patients and inclusion of primary progressive HD. In addition, multivariate analysis were not performed systematically.7,8,9

Brice et al performed one of the largest studies evaluating prognostic factors in relapsed HD. One-hundred and eighty seven patients who relapsed after a first complete remission were included. At first relapse, treatment was conventional (chemo- and/or radiotherapy) in 44% and HDCT followed by ASCT in 56%. Two prognostic factors were identified by multivariate analysis as correlating with both FF2F and OS. These factors were the initial duration of first remission (i.e. < 12 months or > 12 months; \( p < 0.0001 \) and stage at relapse (I-II vs. III-IV); \( p = 0.0013 \)). FF2F was 62% and 32%, respectively, OS was 44% and 87% according to the presence of 0 or 2 parameters, respectively. Laboratory data were not available in this retrospective analysis.10

The GHSG has recently performed a retrospective analysis including a much larger number of relapsed patients \( n=422 \) than previous reported (Figure 1). The analysis of prognostic factors suggests that the prognosis of a patient with relapsed HD can be estimated according to several factors. The most relevant factors were combined into a prognostic score. This score was calculated on the basis of duration of first remission, stage at relapse and the presence or absence of anemia at relapse. Early recurrence within 3 to 12 months after the end of primary treatment, relapse stage III or IV and haemoglobin <10.5g/dl in female or <12g/dl in male patients contribute to a score with possible values 0, 1, 2 and 3 in order of worsening prognosis.11 This prognostic score allows distinguishing patients with different FF2F and OS. The actuarial 4-year FF2F and OS for patients relapsing after chemotherapy with three unfavorable factors were 17% and 27%, respectively. In contrast, patients with none of the unfavorable factors had FF2F and OS at 4-year of 48% and 83%, respec-
tively. In addition, the prognostic score was also predictive for patients relapsing after radiotherapy, for patients relapsing after chemotherapy who were treated with conventional therapies or with HDCT followed by ASCT, and for patients under 60 years and a Karnofsky performance status ≥ 90% being the major candidate groups for dose intensification. Our prognostic factor score uses clinical characteristics which can be easily collected at the time of relapse. It separates groups of patients with substantially different outcomes.

The prognostic factors identified may be useful to tailor the therapy for subgroups of patients, to define homogeneous cohorts for prospective randomized trials, and to identify more precisely patients with poor-risk relapse who should be treated with innovative approaches.

**Treatment strategies**

The survival of patients treated with conventional chemotherapy after relapse of irradiated early-stage disease is at least equal to that of advanced-stage patients initially treated with chemotherapy. Overall survival (OS) and disease-free survival (DFS) range from 57% to 71%. Patients who relapse following radiation therapy alone for localized Hodgkin’s disease have satisfactory results with combination chemotherapy and are not considered candidates for HDCT and ASCT.

HDCT followed by ASCT has been shown to produce 30%-65% long-term disease-free survival in selected patients with refractory and relapsed HD. In addition, the reduction of early transplant-related mortality from 10% - 25% reported in earlier studies to less than 5% in more recent studies has led to the widespread acceptance of HDCT and ASCT.

Although results of HDCT have generally been better than those observed after conventional-dose salvage therapy, the validity of these results has been questioned due to the lack of randomized trials. The most compelling evidence for the superiority of HDCT and ASCT in relapsed HD comes from two reports from the British National Lymphoma Investigation (BNLI) and the German Hodgkin’s Lymphoma Study Group (GHSG) together with the European Group for Blood and Marrow Transplantation (EBMT).

In the BNLI trial, patients with relapsed or refractory HD were treated with a combination of carmustine (BCNU), etoposide, cytarabine and melphalan at a conventional-dose level (mini-BEAM) or a high-dose level (BEAM) with autologous bone-marrow transplantation. The actuarial 3-year event-free survival (EFS) was significantly better in patients who received high-dose chemotherapy (53% vs 10%).

The largest randomized, multicenter trial was performed by the GHSG/EBMT to determine the benefit of HDCT in relapsed HD. Patients with relapse after polychemotherapy were randomly assigned between four cycles of Dexa-BEAM (dexamethasone, BCNU, etoposide, Ara-C and melphalan) and two cycles of Dexa-BEAM followed by HDCT (BEAM) and ABMT/PBSCT. The final analysis of 144 evaluable patients revealed that from 117 patients with PR or CR after two cycles of chemotherapy, FFTF in the HDCT group was 55% versus 34% for the patients receiving an additional two cycles of chemotherapy. OS was not significantly different.

**Sequential high-dose chemotherapy**

In recent years, sequential high-dose chemotherapy has increasingly been employed in the treatment of solid tumors, hematologic and lymphoproliferative disorders. Initial results from phase-I/II studies indicate that this kind of therapy offers safe and effective treatment. In accordance with the Norton-Simon hypothesis, following initial cytoreduction, few non-cross-resistant agents are given at short intervals. In general, the transplantation of PBSC and the use of growth factors allow the application of the most effective drugs at the highest possible doses at intervals of one to three weeks. Sequential high-dose chemotherapy thereby enables the highest possible dosing over a minimum period of time (dose intensification).

In 1997 a multicenter phase-II trial with a high-dose sequential chemotherapy program and a final myeloablative course was started to evaluate the feasibility and efficacy of this novel regimen in patients with relapsed HD. Eligibility criteria included age 18-60 yrs., histologically proven relapsed or primary progressive HD, second relapse with no prior HDCT and ECOG performance status 0-1.

The treatment program consists of two cycles of DHAP (dexamethasone, ara-C, cisplatin) in the first phase in order to reduce tumor burden before HDCT. Patients with partial remission (PR) or complete remission (CR) after two cycles of DHAP,
receive sequential high-dose chemotherapy consisting of cyclophosphamide 4 g/m² iv, methotrexate 8 g/m² iv plus vincristine 1.4 mg/m² iv; and etoposide 2 g/m² iv. The final myeloablative course was BEAM followed by PBSCT with at least 2 x 10⁶ CD34+ cells/kg.

At the last interim analysis 102 patients were available for the final evaluation. State of remission was multiple relapse in 10 patient, progressive disease in 16 patients and late relapse in 44 patients. At 18 months of median follow-up (range 3-31 months) results are as follows: Response rate (RR) after DHAP 87% (23% CR, 64% PR) and RR at final evaluation 77% (68% CR, 9% PR). Toxicity was tolerable with no treatment related deaths. FFTF and OS for patients with early relapse were 64%/87% for early relapse; 68%/81% for late relapse; 30%/58% for patients with progressive disease and 55%/88% for patients with multiple relapse.

In conclusion, sequential administration of high doses of cyclophosphamide, methotrexate and etoposide is feasible and did not affect the tolerability of final myeloablative BEAM. This new, three-phase treatment regimen is well tolerated and feasible in patients with relapsed and primary progressive HD. The preliminary data suggests a high efficacy in relapsed HD patients, warranting further randomized studies.

**HDR-2 Protocol**

In January 2001, the GHSG together with the EORTC, the GEL/TAMO and the EBMT started a prospective randomized study to compare the effectiveness of a standard HDCT (BEAM) with a sequential HDCT after initial cytoreduction with 2 cycles DHAP (HD-R2 protocol, Fig. 2).

Patients with histologically confirmed early or late relapsed HD, and patients in second relapse with no prior HDCT fulfilling the entry criteria receive two cycles of dexamethasone, high-dose cytarabine and cisplatin (DHAP) followed by G-CSF.

Patients achieving NC, PR or CR after DHAP are centrally randomized to receive either BEAM followed by PBSCT (arm A of the study) or HD cyclophosphamide + G-CSF, followed by HD-MTX + vincristine, followed by HD etoposide + G-CSF and a final myeloablative course with BEAM (arm B of the study).

**Allogeneic transplantation after reduced conditioning in HD**

Allogeneic transplantation (alloBMT) has clear advantages compared with autologous transplantation: Donor marrow cells uninvolved by malignancy are used avoiding the risk of infusing occult lymphoma cells, which may contribute to relapse in patients who undergo autologous transplantation. In addition, donor lymphoid cells can potentially mediate a graft-versus-lymphoma effect.

Generally, donor availability and age constraints have limited a broader application of alloBMT in HD. Moreover, alloBMT is associated with a high treatment related mortality rate of up to 75% observed in patients with induction failure which casts doubt upon the feasibility of this approach in HD patients. In most cases, allogeneic transplantation from HLA-identical siblings is not recommended for patients with HD. The reduced relapse-rate associated with a potential graft-versus-tumor effect is offset by lethal graft-versus-host toxicity.

Nevertheless, patients with induction failure and relapsed patients with additional risk factors have a poor prognosis also after HDCT and ASCT. Therefore, the role of alloBMT should be further evaluated in these patients taken advantage of new developments like non-myeloablative conditioning regimens and alloPBSCT.

To circumvent the problems inherent to the toxicity and treatment related mortality associated to allografting, the possibility to achieve engraftment of allogeneic stem cells after immunosuppressive therapy combined with myelosuppressive but non-myeloablative therapy has been assessed. Several groups have recently updated their experience with non-myeloablative conditioning regimens.

The EBMT together with the GEL/TAMO, the EORTC and the GHSG activated a multicenter phase II study to evaluate the treatment-related mortality (TRM) of patients with primary progressive or relapsed HD (early relapse, multiple relapse and relapse after autologous SCT). Patients with an HLA compatible sibling donor or an HLA matched unrelated donor will be initially treated with 1-2 cycles of DHAP or other salvage protocols to reduce tumor burden before alloPBSCT. PBSC will be collected after G-CSF priming of the donor and reinfused after conditioning with fludarabine and melphalan.

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Future directions

Alternative strategies have been developed to improve the outcome of relapsed and resistant HD. These approaches include the development of new cytostatic drugs and biological agents with proven efficacy in preclinical models.

One of the most promising new cytostatic drugs is the new vinca alkaloid vinorelbine, which has demonstrated activity in HD even in patients pretreated with vincristine or vinblastine.34 The use of vinorelbine in first and second-line therapy of HD in order to improve frequency and duration of response is still under investigation. The pyrimidine analogue gemcitabine is the only drug currently under investigation that represents a new cytostatic mechanism of action. The "self-potentiating" mechanism of action leads to an enhanced accumulation and prolonged retention of gemcitabine in the malignant cell. The results of gemcitabine in advanced relapsed HD are promising, with an overall response rate of 53% in heavily pretreated patients.35

Although some clinical efficacy has been demonstrated in clinical trials with immunotoxins (IT) none of the current available IT seems to be suited for a clinical phase-III study.36-38 Bispecific monoclonal antibodies (BiMoab) such as the recently reported CD30xCD64 BiMoab look more promising with clinical development programs scheduled including phase III. The use of recombinant DNA technology for site-directed modifications of the IT and the development of humanized IT and BiMoabs might optimize their efficacy.38 In the future, combining standard chemo-/radiotherapy with biological agents might result in the elimination of residual tumor cells and subsequently more relapse-free long term survivors.

References
9. Reece D, Barnett M, Shepherd J, et al: High-dose cyclophosphamide, carmustine (BCNU), and etoposide (VP16-213) with or without cisplatin (CBV +/- P) and autologous transplantation for patients with Hodgkin’s disease who fail to enter a complete remission after combination chemotherapy. Blood 1995, 86, 451-458


**Introduction**

In the 19th century chronic myeloid leukemia (CML) and polycythemia vera (PV) have been described as primary distinct disease entities. In 1960 Nowell and Hungerford described the presence of a minute chromosome in leukemic cells of patients with chronic myeloid leukemia (CML). This minute chromosome was called Philadelphia chromosome or Ph after the city of discovery. Using banding techniques Janet Rowley (1973) discovered that the Ph originated from a translocation between the long arms of chromosomes 9 and 22, 9(9::22)(q34;q11). Two groups working together in scientific friendship discovered that a hybrid gene is generated by the translocation consisting of the BCR gene on chromosome 22 and the ABL oncogene originating from chromosome 9. This results in a BCR/ABL fusion gene with high tyrosine kinase activity and CML-transformation capacity. Ninety-five percent of all CML patients are Ph+; 90% are Ph+/BCR/ABL+, 5% are Ph+/BCR/ABL-, and 5% are Ph-BCR/ABL-, the latter group usually diagnosed as atypical CML, juvenile CML, chronic neutrophilic leukemia or chronic myelomonocytic leukemia. According to strict morphological, biochemical, cytogenetic and molecular criteria including the Philadelphia (Ph) chromosome and bcr/abl fusion gene and protein, chronic myeloid leukemia is a malignant disease with an obligate transition into acute leukemia, whereas essential thrombocythemia (ET), polycythemia vera (PV) and agnogenic myeloid metaplasia (AMM) form the Ph-chromosome and bcr/abl negative chronic myeloproliferative disorder (MPD) featured by a benign proliferation of the 3 hematopoietic cell lines. Increase and clustering of abnormal enlarged megakaryocytes together with no or various degrees of increased erythropoiesis and/or granulopoiesis in bone marrow biopsy specimens appears to be a pathognomonic clue to the diagnosis of pre-fibrotic MPD when WHO bone marrow features are applied. On top of established WHO bone marrow features in combination with JAK2V617F mutation screening, and laboratory features including endogenous erythroid colony (EEC) formation and serum erythropoietin (EPO) levels we propose updated European clinical, molecular and pathological (ECMP) criteria for the differential diagnosis of true ET, PV and chronic idiopathic myelofibrosis (CIMF) or AMM. The ECMP criteria reduced the platelet count of 600 x 10^9/l to the upper limit of normal (>400 x 10^9/l) as inclusion criterion for the diagnosis of thrombocythemia in various MPDs. When WHO and ECMP criteria are applied, PVSG-defined ET includes three distinct entities; true ET, early PV mimicking ET, and thrombocythemia associated with early CIMF-0 mimicking ET. Compared to characteristic bone marrow features pathognomonic for MPD, spontaneous EEC and low serum EPO levels are not sensitive enough as isolated markers for the diagnosis and differential diagnosis of pre-fibrotic ET, PV and CIMF-0. Bone marrow histology assessment remains the gold standard criterion for the diagnosis and staging of the MPDs. PVSG-defined ET and the presence of the JAK2V617F mutation, EEC, PRV-1 overexpression and low serum EPO levels is consistent with early PV ("forme fruste" PV) when ECMP criteria are applied. The combination of JAK2V617F mutation and increased hematocrit (>0.051 males and >0.48 females) is consistent with the diagnosis PV (specificity 100%, sensitivity 95%) without the need of red cell mass measurement. About half of the ET and CIMF patients are JAK2V617F positive (sensitivity 50%). The degree of JAK2V617F positivity of granulocytes is related to disease stage: heterozygous in true ET and early PV and mixed hetero/homozygous to homozygous in overt and advanced PV and CIMF-1 to 3. The proposed ECMP criteria for the differential diagnosis of ET, PV, CIMF in patients with JAK2V617F positive and JAK2 wild type MPD should be evaluated in prospective management studies in search for the most relevant prognostic factors of therapeutic significance.

**Key words:** myeloproliferative disorders, essential thrombocythemia, polycythemia vera, chronic idiopathic myelofibrosis, erythropoietin, endogenous erythroid colony assay, JAK2 V617F mutation, bone marrow pathology.
myeloid metaplasia (AMM) form the Ph-chromosome and brc/abl negative chronic myeloproliferative disorder featured by a benign proliferation of the 3 hematopoietic cell lines (figure 1)\(^{10}\).

In 1950, William Dameshek described polycythemia vera (PV) as a chronic disorder of the bone marrow characterized by excessive production of nucleated red cells, granulocytes and megakaryocytes peripheral blood erythrocytosis, leukocytosis and thrombocytosis (figure 1)\(^{11}\). Some cases however show only a moderate elevation of erythrocytes with an extreme degree of thrombocytosis, while in others the leukocyte counts may be at or close to leukemic levels, with only slight increase in red cells or platelets\(^{11}\). According to Dameshek all “stops” to blood production in the bone marrow seem to have been pulled out in PV. As to the etiology of PV, Dameshek proposed two highly speculative possibilities: first, the presence of excessive bone marrow stimulation by an unknown factor or factors, and second, a lack or a diminution in the normal inhibitory factor or factors\(^{10}\). This hypothesis of Dameshek has recently confirmed by the discovery of the JAK2V617F mutation\(^{12}\) demonstrating that the V617F mutation induces a loss of inhibitory activity of the JH2 pseudokinase part on the JH1 kinase part of JAK2, leading to enhanced activity of the normal JH1 kinase activity of JAK2, which makes the mutated hematopoietic stem cells hypersensitive to hematopoietic growth factors TPO, EPO, IGF1, SCF and GCSF, resulting in trilinear myeloproliferation (Figure 1)\(^{13}\).  

### The concept of PV as a trilinear MPD

Wasserman extended in 1954 the 1950 concept of Dameshek on PV as a trilinear MPD and distinguished at least five subsequent stages in the natural history of PV\(^{3}\). 

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### Table 1. Polycythemia Vera Study Group (PVSG) criteria for the diagnosis of essential thrombocythemia (ET), polycythemia vera (PV) and modified criteria for the diagnosis of chronic idiopathic myelofibrosis (CIMF)

<table>
<thead>
<tr>
<th>PVSG criteria for the diagnosis of ET17,18</th>
<th>Pathological criteria (WHO)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Platelet count &gt; 500 x 10^9/L (ECMP &gt; 400 x 10^9/L)</td>
<td>C1. Increase of dispersed or loosely clustered, predominantly enlarged megakaryocytes with mature cytoplasm and hyperlobulated nuclei (figures 2 and 3)</td>
</tr>
<tr>
<td>- No known cause of Reactive Thrombocytosis</td>
<td>P1. Increase of dispersed or loosely clustered megakaryocytes with mature cytoplasm and hyperlobulated nuclei</td>
</tr>
<tr>
<td>- Normal hemoglobin and red cell mass to exclude overt PV</td>
<td>C2. Presence of large or giant platelets in a peripheral blood smear</td>
</tr>
<tr>
<td>- Stainable iron bone marrow to exclude PV</td>
<td>C3. Presence of MPL(^{30}) or Jak2V617F mutation</td>
</tr>
<tr>
<td>- No tear drop erythrocytes and no leuko-erythroblastosis</td>
<td>C4. Normal values for hemoglobin, hematocrit, white blood cell differential count</td>
</tr>
<tr>
<td>- No features of MDS in bone marrow smear and biopsy</td>
<td>C5. Absence of the Philadelphia chromosome or any other cytogenetic fusion-gene abnormality</td>
</tr>
<tr>
<td>- Absence of Ph + chromosome (brc/abl) to exclude CML</td>
<td></td>
</tr>
<tr>
<td>- Myelofibrosis grade 1 up to grade 2 is allowed in case of collagen fibrosis of bone marrow is allowed up to &lt;1/3 of bone marrow biopsy area.</td>
<td></td>
</tr>
</tbody>
</table>

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### Table 2. WHO bone marrow features and European clinical, molecular and pathological (ECMP) criteria for the diagnosis of essential thrombocythemia (true ET)\(^{14,15}\)

<table>
<thead>
<tr>
<th>Clinical and molecular criteria</th>
<th>Pathological criteria (WHO)</th>
</tr>
</thead>
<tbody>
<tr>
<td>A. Major criteria</td>
<td>C1. Sustained platelet count above the upper limit of normal: &gt;400 x10^9/L</td>
</tr>
<tr>
<td>B. Minor criteria</td>
<td>C2. Presence of large or giant platelets in a peripheral blood smear</td>
</tr>
<tr>
<td>C. Minor criteria</td>
<td>C3. Presence of MPL(^{30}) or Jak2V617F mutation</td>
</tr>
<tr>
<td>D. Minor criteria</td>
<td>C4. Normal values for hemoglobin, hematocrit, white blood cell differential count</td>
</tr>
<tr>
<td>E. Minor criteria</td>
<td>C5. Absence of the Philadelphia chromosome or any other cytogenetic fusion-gene abnormality</td>
</tr>
</tbody>
</table>

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Figure 1. The concept of Dameshek in 1950 on polycythemia vera (PV) as a trilinear myeloproliferative disorder (MPD) due to one hypothetical stimulus, appeared to be caused by the acquired JAK2V617F mutation discovered by Vainchenker et al in 2005. The unifying concept of Dameshek in 1951 on the chronic myeloproliferative disorders (CMPDs) essential thrombocythemia (ET), polycythemia vera (PV), agnogenic myeloid metaplasia (AMM) has been broken up by the PVSG into Ph-positive MPDs and CML complicated by myelofibrosis (MF) and the Ph-negative MPDs ET, PV and MF either positive or negative for the acquired JAK2V617F mutation.
Treated by thrombocythemia, erythrocythemia and size, which is labelled as idiopathic erythrocytosis.

Increased hemoglobin, hematocrit and red cell mass with normal leukocytes, thrombocytes and spleen.

CIMF-0 or CIMF-1

Normal or decreased platelet count

Leukocytosis, leukopenia

Pronounced splenomegaly

Grading of myelofibrosis (MF):

MF 1: early fibrotic CIMF-1: loose network of reticulin with many intersections, especially in peripheral areas, no collagenization

MF 2: fibrotic CIMF-2: diffuse and dense increase in reticulin with extensive intersections, occasionally with only focal bundles of collagen and/or focal osteosclerosis.

MF 3: classic CIMF-3: diffuse and dense increased in reticulin with extensive interactions with coarse bundles of collagen often associated with significant osteosclerosis

MF > 3: endstage hypocellular with extensive osteomyelosclerosis

Table 4. WHO bone marrow features and European clinical, molecular and pathological (ECMP) Criteria for the Diagnosis of Polycythemia Vera (PV) and diagnostic differentiation between PV and congenital or acquired erythrocytosis

Clinical and molecular criteria

Pathological criteria (WHO)

Major

A 0. Early PV. Hematocrit in the upper limit of normal: Ht: 0.45 to 0.5 in male and 0.43 to 0.48 in female

A 1. Classical PV: Hematocrit > 0.51/0.48 in male/female

A 2. Presence of JAK2V617F mutation

A 3. Low serum Epo level

Minor

B 1. Persistent increase of platelet count: grade I: 400-1500; grade II: >1500.

B 2. Granulocytes >10 x109/l or Leukocytes > 12 x109/l and/or raised LAP-score or increased PRV-1 expression in the absence of fever or infection

B 3. Splenomegaly on palpation or on ultrasonic echogram (>12 cm length in diameter).

B 4. Spontaneous endogenous erythroid colony (EEC) formation (optional)

WHO/ECMP criteria for early and overt PV

A0, A2, B1 and P1 establish early PV mimicking ET PV ECP stage 0, or masked PV

A1, A2, P1 and one or more of B establish polycythemic PV ECP stage 1

A1, A2, P1 and one or more of B establish advanced PV ECP stage 2 and 3

P1 and P2 with normal or increased values of serum EPO is consistent with erythrocytosis

A3 confirms PV. B4 is an important research option.

According to WHO/ECMP criteria, C 1 and P1 plus P2 establish CIMF – any other peripheral blood criteria and grading of secondary myelofibrosis (MF) contribute to ECMP staging of WHO defined CIMF.

Stage 1). Pure erythrocytosis is featured by increased hemoglobin, hematocrit and red cell mass with normal leukocytes, thrombocytes and spleen size, which is labelled as idiopathic erythrocytosis.

Stage 2). The polycythemic stage of PV is featured by thrombocytoma, erythrocytoma and no or slight myeloid metaplasia, leukocytosis and/or splenomegaly.

Stage 3). Myeloid metaplasia in PV patients presents with no or different grades of reticulin and collagen fibrosis in the bone marrow and progressive splenomegaly during long-term follow in about one third of the cases.

Stage 4). The polycythemic stage with various degrees myelofibrosis and splenomegaly following PV may elapse 5 to 25 years before a period of normal red cell values so-called spontaneous remission of PV occurs. This stage must be considered as the beginning of spent phase PV and may last a few to several years. At this point the spleen is frequently large and very firm to palpation, the liver is enlarged to a moderately degree in most patients, thrombocythemia is frequent and may be pronounced with bizarre and giant platelets, and white cells are usually increased with granulocytic leukocytosis (leukocytosis) accompanied by a small percentage of immature forms.

Stage 5). Post-PV myeloid metaplasia shows various degrees of leuko-erythroblastosis of the peripheral blood and may progress to extreme myelofibrosis with a dry tap on aspiration and massive splenomegaly. At this end-stage histopathology of bone marrow biopsy shows a similar picture and
can not been differentiated from agnogenic myeloid metaplasia with no previous history of PV.

**Three phenotypes of MPD: ET, PV and CIMF**

The criteria the Polycythemia Vera Study Group (PVSG) originate from the early 1970s to classify the chronic myeloproliferative disorders (MPD) as 3 disease entities ET, PV and idiopathic myelofibrosis (IMF) and separate these 3 MPDs from Philadelphia chromosome-positive chronic myeloid leukaemia (CML) (figure 1, table 1)\(^1\). The criteria of Ph-chromosome-negative myeloproliferative disorders according to the 1990 Hannover bone marrow classification\(^9\), the Rotterdam clinical and pathological criteria between 1997 and 2000\(^2\) and the World Health Organisation (WHO) in 2001\(^2\) are an attempt to integrate bone marrow morphological criteria alongside PVSG clinical criteria for essential thrombocytthemia (ET), polycythemia vera (PV) and chronic idiopathic myelofibrosis (CIMF). However, early stages of PV and ET are not recognized by the PVSG and 2001 WHO classifications and many patients will be diagnosed as unclassifiable MPD, indicating the need to modify the diagnostic inclusion criteria for early and overt stages of MPD. In the present study we propose a new set of European clinical, molecular and pathological (ECMP) criteria in a joint effort by clinicians and pathologists to integrate PVSG criteria, WHO bone marrow features and the use of new laboratory and molecular markers for diagnostic differentiation of each of the early and overt MPD phenotypes more accurately as a sound basis on which proper treatment guidelines are to be recommended.

---

**Table 5. ET according to PVSG\(^1\), WHO\(^2\) and ECMP\(^2\) criteria**

<table>
<thead>
<tr>
<th>ECMP PVSG</th>
<th>Hereditary ET</th>
<th>True ET type 1</th>
<th>Early PV ET type 2</th>
<th>Proibiotic CIMF ET type 3</th>
</tr>
</thead>
<tbody>
<tr>
<td>Incidence</td>
<td>&lt; 0.001</td>
<td>20-30</td>
<td>20-30</td>
<td>40-60</td>
</tr>
<tr>
<td>Serum EPO</td>
<td>Normal</td>
<td>Normal</td>
<td>Decreased</td>
<td>Normal</td>
</tr>
<tr>
<td>Platelets</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Erythrocytes</td>
<td>N</td>
<td>N</td>
<td>N/↑</td>
<td>N/↓</td>
</tr>
<tr>
<td>Hematocrit</td>
<td>N</td>
<td>N</td>
<td>N/↑</td>
<td>N/↓</td>
</tr>
<tr>
<td>Bone marrow: ET picture</td>
<td>ET picture</td>
<td>ET picture</td>
<td>PV picture</td>
<td>CIMF picture</td>
</tr>
<tr>
<td>Megakaryocytes</td>
<td>Normal large / giant and mature</td>
<td>Abnormal</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Splenomegaly</td>
<td>-</td>
<td>/ +</td>
<td>-/ +</td>
<td>+ / -</td>
</tr>
<tr>
<td>JAK2 V617F</td>
<td>Neg:</td>
<td>-</td>
<td>+/ +</td>
<td>+ / -</td>
</tr>
<tr>
<td>EEC</td>
<td>-</td>
<td>/ +</td>
<td>++</td>
<td>+ / -</td>
</tr>
<tr>
<td>PVR-1</td>
<td>-</td>
<td>/ +</td>
<td>++</td>
<td>+ / -</td>
</tr>
<tr>
<td>Clonality</td>
<td>polyclonal</td>
<td>monoclonal</td>
<td>Monoclonal</td>
<td>Monoclonal</td>
</tr>
</tbody>
</table>

---

**Table 6. Staging of PV patients according to WHO/ECMP criteria: therapeutic implications\(^2\)**

<table>
<thead>
<tr>
<th>PV, ECP stage</th>
<th>0</th>
<th>1</th>
<th>2</th>
<th>3</th>
<th>4</th>
<th>5</th>
</tr>
</thead>
<tbody>
<tr>
<td>Clinical Diagnosis</td>
<td>Early PV ET type 2</td>
<td>polycythemic PV</td>
<td>Classic PV</td>
<td>Advanced PV</td>
<td>Post-PV myelofibrosis</td>
<td>Spent phase PV</td>
</tr>
<tr>
<td>LMP-score and/or PVR-1</td>
<td>↑</td>
<td>↑</td>
<td>↑</td>
<td>↑/↑↑↑</td>
<td>variable</td>
<td>variable</td>
</tr>
<tr>
<td>Red cell mass</td>
<td>N</td>
<td>↑</td>
<td>↑</td>
<td>↑</td>
<td>variable</td>
<td>N/↓</td>
</tr>
<tr>
<td>Serum EPO</td>
<td>N/↓</td>
<td>↓</td>
<td>↓</td>
<td>↓</td>
<td>variable</td>
<td>N/↓</td>
</tr>
<tr>
<td>Leukocytes x10⁹/l</td>
<td>&lt; 12</td>
<td>&lt; 12</td>
<td>N-&gt;12</td>
<td>&gt; 15</td>
<td>&gt; 20</td>
<td>&gt; 20</td>
</tr>
<tr>
<td>Platelets x10⁹/l</td>
<td>&gt; 400</td>
<td>&lt; 400</td>
<td>&gt; 400</td>
<td>&gt; 1000</td>
<td>variable</td>
<td>variable</td>
</tr>
<tr>
<td>Periperal blood red cells</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Hemoglobin g/dl (mmol/l)</td>
<td>&lt; 12(10)</td>
<td>&gt; 16(10)</td>
<td>&gt; 16(10)</td>
<td>&gt; 16(10)</td>
<td>variable</td>
<td>N/↓</td>
</tr>
<tr>
<td>Hematocrit</td>
<td>&lt; 0.51</td>
<td>&gt; 0.51</td>
<td>&gt; 0.51</td>
<td>&gt; 0.51</td>
<td>variable</td>
<td>N/↓</td>
</tr>
<tr>
<td>Erythrocytes x10¹²/l</td>
<td>&lt; 6</td>
<td>&gt; 6</td>
<td>&gt; 6</td>
<td>&gt; 6</td>
<td>variable</td>
<td>N/↓</td>
</tr>
<tr>
<td>WHO bone marrow</td>
<td>Early PV</td>
<td>Early PV</td>
<td>Trilinear PV</td>
<td>Trilinear PV</td>
<td>Trilinear /MF</td>
<td>MF</td>
</tr>
<tr>
<td>Bone marrow cellularity (%)</td>
<td>50-80</td>
<td>50-80</td>
<td>80-100</td>
<td>80-100</td>
<td>Decreased</td>
<td>Decreased</td>
</tr>
<tr>
<td>Grading myelofibrosis</td>
<td>MF 0</td>
<td>MF 0</td>
<td>MF 0</td>
<td>MF 0</td>
<td>MF 0/1</td>
<td>MF 3</td>
</tr>
<tr>
<td>Splenomegaly</td>
<td>slight</td>
<td>no</td>
<td>Slight/moderate</td>
<td>moderate</td>
<td>large</td>
<td>large</td>
</tr>
<tr>
<td>Spleen size, echogram cm</td>
<td>12-15</td>
<td>12-15</td>
<td>12-15</td>
<td>12-20</td>
<td>&gt; 20</td>
<td>&gt; 20</td>
</tr>
<tr>
<td>Specific MPD markers</td>
<td>EEC</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+/↑?</td>
</tr>
<tr>
<td>Molecular JAK2V617F</td>
<td>+</td>
<td>+</td>
<td>+/++</td>
<td>+/++</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>Granulocytes</td>
<td>+</td>
<td>+</td>
<td>+/++</td>
<td>+/++</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>BFU-e</td>
<td>++</td>
<td>++</td>
<td>++</td>
<td>++</td>
<td>++</td>
<td>++</td>
</tr>
<tr>
<td>Therapeutic implications</td>
<td>First line treatment</td>
<td>Aspirin</td>
<td>Phlebotomy aspin</td>
<td>Phlebotomy* aspin</td>
<td>INF/HU* aspin</td>
<td>HU</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Supportive Splenectomy</td>
<td></td>
</tr>
</tbody>
</table>

*↑ = increased, ↓ = decreased, N = normal, + = present or heterozygous; ++ = homozygous
Limitations of PVSG and 2001 WHO criteria for the diagnoses of ET, PV and CIMF

The Polycythemia Vera Study Group (PVSG) reduced since 1986 the platelet count of 1000 to 600 x10⁹/l as the arbitrary minimum for the diagnosis of ET (table 1)¹⁷,¹⁸. Lengfelder et al showed that this minimum of 600 x10⁹/l platelets according to the PVSG excluded early stage ET at platelets between normal and 600 x10⁹/l in 29% of 143 ET cases when bone marrow biopsy was included in the investigations employed to diagnose ET²². These data confirmed the need to modify the PVSG criteria by lowering the platelet counts to 400 x10⁹/l as the upper limit of normal for the clinical diagnosis of ET (table 2)²⁰,²³,²⁴.

The 2001 WHO criteria combined a typical ET histological bone marrow picture with platelet count in excess of 600 x10⁹/l for the diagnosis of ET²¹, thereby excluding early stage ET²²-²⁴. Only one-third of PVSG-defined ET is diagnosed as true ET when the 2001 WHO clinical and bone marrow criteria are applied²⁵. Unclassifiable early stage CMPD according to the 2001 WHO criteria include 3 types of early stage MPD: 1) initial ET with a typical ET bone marrow but platelet count below 600 x10⁹/l; 2) initial PV with a typical PV bone marrow, platelet count less than 600 x10⁹/l, low serum erythropoietin (EPO), normal red cell mass and hematocrit less than 0.51; and 3) initial masked MPD with splenomegaly and normal or slightly increased platelet count and hematocrit²¹.

According to Wasserman, PV frequently presents with initial erythrocytosis (idiopathic erythrocytosis), and distinguished at least five subsequent stages in the natural history of PV³. The PVSG used 3 major and 4 minor clinical criteria as inclusion criteria for the diagnosis of PV in the PVSG-01 study (Table 1)¹⁶,¹⁷. Increased red cell mass is a crude inclusion criterion for the diagnosis of PV (Louis Wasserman personal communication ASH 1995) and corresponds to high hematocrit values between 0.48 and 0.76, increased platelet count above 400 x10⁹/L in two-third and palpable spleen in two-third of about 400 PV patients in the PVSG-

---

**Table 7. Translation of PVSG defined ET into WHO-defined True ET and Chronic Idiopathic Myelofibrosis (CIMF) grade 0, 1, 2, and 3 in a retrospective study of 865 Thrombocythemia patients²⁰,²¹,²².**

<table>
<thead>
<tr>
<th>PVSG clinical diagnosis</th>
<th>WHO pathological diagnosis</th>
<th>ET</th>
<th>True ET</th>
<th>ET</th>
<th>CIMF-0</th>
<th>ET</th>
<th>CIMF-1</th>
<th>Classic IMF</th>
<th>CIMF-2</th>
<th>Classic IMF</th>
<th>CIMF-3</th>
</tr>
</thead>
<tbody>
<tr>
<td>WHO bone marrow</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ec Grading MF³⁰</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Number of evaluable patients</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Age years (mean)</td>
<td>59</td>
<td>67</td>
<td>69</td>
<td>70</td>
<td>-</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Platelets (mean x10⁹/l)</td>
<td>1176</td>
<td>902</td>
<td>841</td>
<td>649</td>
<td>308</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Leukocytes (mean x10⁹/l)</td>
<td>11.1</td>
<td>12.4</td>
<td>12.2</td>
<td>11.0</td>
<td>10.9</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Anemia</td>
<td>No</td>
<td>No</td>
<td>No</td>
<td>moderate</td>
<td>Moderate/severe</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Hemoglobin: g/dl (mean)</td>
<td>13.8</td>
<td>13.9</td>
<td>13.1</td>
<td>12.0</td>
<td>10.6</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>mmol/l (mean)</td>
<td>8.6</td>
<td>8.7</td>
<td>8.2</td>
<td>7.5</td>
<td>6.6</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Leuko-erythroblastosis and tear drop erythrocytes:</td>
<td>No</td>
<td>No</td>
<td>No/slight</td>
<td>Clearly present</td>
<td>Pronounced present</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Splenomegaly*</td>
<td>No/slight</td>
<td>no/slight</td>
<td>No*/slight</td>
<td>Moderate</td>
<td>Pronounced</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Palpable spleen cm*</td>
<td>0.4</td>
<td>1.1</td>
<td>1.4</td>
<td>3.2</td>
<td>5.5</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>LDH increase above the upper limit of normal</td>
<td>no</td>
<td>no</td>
<td>no*/slight</td>
<td>yes</td>
<td>yes</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Adverse signs**</td>
<td>no</td>
<td>no</td>
<td>no</td>
<td>no</td>
<td>1 or 2</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mean life expectancy, years</td>
<td>&gt; 20</td>
<td>&gt; 15</td>
<td>&gt; 15 to &gt; 10</td>
<td>Around 10</td>
<td>&lt;5</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

* Spleen and liver size on palpation in cm below the costal margin.
** Adverse signs include age > 70 years, hemoglobin < 10 g/dl, myeloblasts PB > 2%, erythro-normoblasts PB > 2%, leukocytosis > 20 x 10⁹/l, thrombocytopenia < 300 x 10⁹/l, severe constitutional symptoms, massive splenomegaly, cytogenetic abnormalities.
Evidence accumulates that the very early prefibrotic stage of chronic myeloproliferative disease is featured by a platelet count in the upper limit of normal (around 400 x10^9/L), the presence of enlarged platelets in a peripheral blood smear in the complete absence of any underlying disorder for reactive thrombocytosis. ET according to the PVSG criteria comprises 3 types of prefibrotic MPDs including true ET, thrombocythemia associated with early polycythemia vera (PV), and thrombocythemia associated with prefibrotic chronic idiopathic myelofibrosis (CIMF-0) when the recently defined WHO bone marrow features and the European clinical molecular and pathological (ECMP) criteria are applied (figure 2, table 5). We produced good evidence that characteristic PV bone marrow histology features (irrespective of red cell mass measurements) are seen in four different stages of newly diagnosed MPD patients (figures 3, tables 4 and 6): First, early PV mimicking ET with a hematocrit in the upper limit of normal but increased platelet count without or with slight splenomegaly (stage 0 PV, table 6); Second, erythrocytosis with increased red cell mass, high hematocrit, low serum erythropoietin (EPO), but normal platelet count and spleen size (so-called idiopathic erythrocytosis = stage 1 PV, table 6); Third, classic PV with increased red cell mass, high hematocrit and one or more PVSG B criteria PV (overt stage 2 and 3 PV, table 6); Fourth, unclassifiable MPD or masked PV with pronounced splenomegaly, normal hemoglobin and hematocrit, normal or slightly elevated platelet count. Thiele et al has nicely worked out this concept and confirmed that characteristic PV bone marrow histopathological features are seen in classic PV and in early PV mimicking ET (tables 4 and 6). The 2001 WHO criteria com-
bined a characteristic PV histological bone marrow picture as a minor criterion with increased red cell mass as a major crude inclusion criterion for the diagnosis of PV, thereby excluding early stage PV mimicking ET. To overcome the shortcomings of the 2001 WHO classification of the MPDs, we here propose the updated European clinical, molecular and pathological (ECMP) for the diagnosis, classification and staging of true ET, PV and CIMF (figures 2, and 3, tables 2, 3 and 4).24-28

**Limitations of laboratory markers for the diagnosis of ET and PV**

Red cell mass measurement (RCM) is cumbersome, time consuming, costly, and not specific for MPD. Increased RCM separates patients with increased erythrocytosis from apparent erythrocytosis (pseudo-polycythemia), but does not distinguish PV from congenital polycythemia (CP) or secondary erythrocytosis (SE). The ECMP separates patients with increased erythrocytosis into patients with polycythemia vera (true polycythemia, figure 2 and 3) as a trilineal MPD and non-clonal polycythemia (erythrocytosis, table 4), either congenital, acquired or idiopathic. In a consecutive cohort of 105 patients with both PV and non-clonal polycythemia as well as ET, in whom diagnostic categories were established based on clinical data, laboratory parameters and bone marrow histology, RCM had a sensitivity of 76% in the diagnosis of PV and a specificity of 79% in distinguishing PV and non-clonal polycythemia. PV patients with RCM may have normal hemoglobin and hematocrit because of associated iron deficiency and/or significant splenomegaly. Bone marrow histopathology has a sensitivity and specificity of near to 100% (gold standard) to differentiate a typical trilineal hypercellular bone marrow with small and enlarged (pleomorphic) megakaryocytes in early and overt PV (table 4) from the presence of isolated increased erythrocytosis and normal megakaryocytes in congenital (primary) polycythemia (CP), congenital erythrocytosis (CE) idiopathic erythrocytosis (IE) or acquired (secondary) erythrocytosis (SE) without the need for red cell mass measurement. In none of the PV patients is the diagnosis from RCM measurement found to be of additional diagnostic value, because all PV patients with increased red cell mass not only show a typical PV bone marrow histology picture but also the presence of one or more specific markers endogenous erythroid colony (EEC), low serum EPO, increased platelet count, and/or splenomegaly for the diagnosis of PV.23,27

Spontaneous EEC and low serum EPO levels are specific confirmative criteria for the diagnosis of PV, but have insufficient diagnostic sensitivity as isolated parameters to differentiate between PV, CP, CE, SE, ET and normal controls. About 50% of PVSG defined ET patients show not
only spontaneous EEC but also increased PRV-1 expression together with low serum EPO levels, indicating that EEC/PRV-1-positive ET comprises a biologically distinct subgroup of ET patients reflecting early PV ("forme fruste" PV) that is at risk for progression to overt PV (tables 5 and 6). Considering the finding of clustered enlarged or giant megakaryocytes according to ECMP criteria as diagnostic for MPD (ET, PV or CIMF) in 46 MPD patients with splenichic thrombosis, the sensitivity of increased red cell mass for the diagnosis of MPD was 63%, of low serum EPO level 52%, of EEC 72%, and of splenomegaly 74%, indicating the superiority of bone marrow histopathology to detect masked, early and overt stages of MPD. In patients with splanchic vein thrombosis (Budd-Chiari syndrome or portal vein thrombosis) without signs of overt MPD, the laboratory markers EEC, PRV-1 expression and low serum EPO are insensitive but the combination of JAK2V617F mutation and bone marrow histology assessment is highly sensitive and specific to diagnose early ET and PV.

The role of JAK2V617F mutation in the pathogenesis and classification of trilinear MPD

The discovery of the JAK2 V617F mutation in 2004 by William Vainchenker and his team in France (JAK2 Vainchenker 617 France) was immediately appreciated as a real evolutionary event, and rapidly confirmed in 2005 by several investigators. JAK2 plays an essential role in hematopoiesis by mediating signals from several hematopoietic cytokines including EPO, TPO IL-3 G-CSF, and GMSCF. The JAK2 V617F mutation makes the mutated hematopoietic progenitor cells hypersensitive to these cytokines, thereby leading to growth advantage of the mutated above the non-mutated normal trilinear hematopoietic cells in the bone marrow (figure 4). The JAK2V617F mutation is detectable in CD34+ hematopoietic bone marrow cells, erythroblasts, in cells of spontaneous EEC, blood platelets and granulocytes. Applying allele-specific polymerase chain reaction (PCR) analysis in PVSG-defined MPD patients, a high frequency of the JAK2V617F mutation of 95% (92-97%) in PV, and a lower frequency of 53% (49-57%) in ET and 52% (44-55%) in idiopathic myelofibrosis (IMF) are described. Only 3 to 4% of ET, 24 to 27% of PV and 6 to 18% of IMF patients are homozygous for the JAK2V617F mutation. Two hypotheses have been proposed by Vainchenker's group of French investigators Delhommeau, James, Pisani, Villeval and Casadevall to explain why three different phenotypes of MPD are caused by the same JAK2V617F mutation: the "dosage" hypothesis and the "additional events" hypothesis (figure 4). According to the dosage hypothesis (based on animal studies and different mutation states of JAK2V617F in MPD patients), the level and duration of JAK2V617F directly contribute to the phenotypic diversity of trilinear MPDs. According to this model (based on animal studies and different mutation states of JAK2V617F in MPD patients), the level and duration of JAK2V617F directly contribute to the phenotypic diversity of trilinear MPDs (figure 4). The hypothesis to explain that the level of kinase activity regulates the disease phenotype of MPD is based on different densities of thrombopoietin (TPO) and EPO receptors (TPOR and EPOR) on hematopoietic progenitor cells and on difference of response of TPOR and EPOR to various levels of JAK2V617F activity. TPOR or MPL is expressed at high levels in megakaryocytic cells where it controls TPO physiologic levels. It is possible that activation of a few TPO receptors by low levels of JAK2V617F (heterozygous) is sufficient to send a signal to megakaryocytic cells. A slight increase in numbers of mutated megakaryocytes and platelets (about 200 x10⁹/l mutated platelets) might be enough to produce platelet-mediated microvascular circulation disturbances. Conversely, EPOR is expressed at low levels on hematopoietic progenitor cells and therefore high levels of JAK2V617F may be required to activate EPOR and generate PV-like phenotype. Sustained high levels of JAK2V617F during long-term follow-up subsequently may lead to a high level activation of EPOR and granulocyte colony stimulating factor receptor (G-CSFR) leading to extramedullary hematopoiesis (splenomegaly) and cytokine mediated secondary myelofibrosis. The degree of JAK2V617F positivity (progression from heterozygous to homozygous in figure 4) is strongly cor-

**Figure 6.** Algorithm for diagnostic work-up of patients with suspected polycythemia vera or erythrocytosis.

<table>
<thead>
<tr>
<th>Hematocrit &gt;0.51 male and &gt;0.48 female</th>
</tr>
</thead>
<tbody>
<tr>
<td>Granulocyte JAK2V617F mutation screening and Serum EPO</td>
</tr>
<tr>
<td>JAK2V617F positive</td>
</tr>
<tr>
<td>Both normal</td>
</tr>
<tr>
<td>Serum EPO increased JAK2V617F negative</td>
</tr>
<tr>
<td>Bone Marrow Biopsy: BMB Serum EPO - Spontaneous EEC</td>
</tr>
<tr>
<td>BMB and EEC (Red cell mass optional)</td>
</tr>
<tr>
<td>Search for congenital or acquired erythrocytosis</td>
</tr>
<tr>
<td>Trilineal MPD ECMP staging PV Therapeutic implications</td>
</tr>
<tr>
<td>Normal Erythrocytosis unlikely</td>
</tr>
<tr>
<td>BMB: Increased erythrocytosis Or EEC + : search for primary or secondary erythrocytosis</td>
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related with (polycythemia rubra vera gene) PRV-1 over expression in granulocytes, with the ability to form spontaneous ECC and with progressive post-PV myelofibrosis55,55,57. Scott et al recently showed that BFU-e colonies are already homozygous for the JAK2V617F mutation in PV patients with a heterozygous pattern of JAK2V617F in their peripheral blood granulocytes58. In contrast, the BFU-E colonies from heterozygous patients with ET did not contain a subpopulation of JAK2V617F homozygous cells58. In a recent study, JAK2V617F was detected in 75% of ET (n=60) and in 97% of PV patients (n=62)59. Allelic ratios exceeding 50% JAK2V617F indicating the presence of granulocytes homozygous for JAK2V617F were found in 70% of PV at diagnosis but never in ET59. Passamonti et al produced good evidence that transition from heterozygosity to homozygosity for the JAK2V617F mutation represents a very important step in the progression from classic PV to post-PV myelofibrosis57.

JAK2V617F may be be dependent not only on the amount of heterozygous and homozygous mutant protein, but also on the various pathway regulating JAK2 activity including MPL, JAK2, STAT-3 signalling pathway53. This has led to the recent discovery of the MPLW515L and MPLW515K mutations as the underlying etiology in some ET and CIMF patients60,61. Pikman et al identified a somatic activating mutation in the transmembrane domain of MPL, the thrombopoietin receptor (TPOR), in 4 of 45 (9%) of JAK2 wild type myelofibrosis patients60. In a mouse model transplant assay, expression of the MPLW515L, but not wild type MPL, resulted in a fully penetrant MPD characterized by marked thrombocytosis and leukocytosis with no evidence of polycythemia. Thrombocytosis with leukocytosis was caused by dual proliferation of megakaryopoiesis and granulopoiesis (myelopoesis) with increased numbers and clustering of atypical dysmorphic megakaryocytes in the bone marrow, myelofibrosis and marked splenomegaly due to extramedullary hemopoiesis consistent with true CIMF60. Pardanani et al screened 1182 PVSG-defined MPD patients (318 ET, 242 PV, and 290 IMF) and 64 controls for MPLW515 mutations regardless of JAK2V617F mutation60. MPL mutations either MPLW515L (n=17) or MPLW515K (n=5) was detected in 20 patients (de novo IMF in 12 = 4%, ET in 4 = 1% and post-ET myelofibrosis in 1, but not in PV and controls61. Six cases carried the MPLW515L and JAK2V617F alleles indicating that these alleles have functional complementation in IMF. These experimental and clinical observations indicate that MPLW515 mutation related myelofibrosis may represent a distinct entity of JAK2 wild type IMF distinct from JAK2V617F trilineal MPD. According to the “additional events” hypothesis, alternative and/or additional molecular abnormalities modify, or precede a homozygous state deferred to by the JAK2V617F mutation alone, combinations carrying the JAK2V617F and MPLW515 mutations61 or other combinations of still unknown mutations (figure 4)63. Mechanisms other than mitotic recombination such as duplication of the mutated allele is observed in a proportion of PV and CIMF patients displaying a gain of 9p, mostly due to trisomy 953. Therefore, there may be an overlap between “dosage” and “additional molecular events” hypotheses. Long-term studies are warranted to delineate the chronology and impact of various putative additional molecular abnormalities especially in cases of progressive disease from JAK2V617F positive true ET to PV and subsequent CIMF and in cases with combined JAK2V617F/MPLW515 mutations or JAK2-wild-type/MPLW515 positive ET and CIMF. Sex appears to be a powerful genetic background modifier in JAK2V617F-positive MPDs as ET is more common in females and PV in males.

**WHO bone marrow features and ECMP criteria for the diagnosis of ET, PV and CIMF**

In 1980s Georgii and Thiele independently defined the pathological features of ET, PV and idiopathic myelofibrosis (IMF) or agnogenic myeloid metaplasia (AMM) as derived from histopathological morphology of bone marrow biopsies10,62. ET is defined by persistent increase of platelets in excess of 400 x10^9/l without the Ph1+ chromosome together with monolinear proliferation of mature enlarged megakaryocytes in the bone marrow with normal cellularity, normal erythropoiesis and normal granulopoiesis (figure 2)10,62. PV is defined as a trilineal proliferation of megakaryopoiesis, erythropoiesis and granulopoiesis in which the erythropoiesis was most prominent together with variable degrees of increased platelets, erythrocytes and granulocytes in the peripheral blood in the absence of the Ph1+ chromosome (figure 2)10,62. Georgii regarded myelofibrosis (MF) as a reactive feature secondary to progressive disease9,10 seen in CMGM, PV and CML19,63,64. Therefore, according to Georgii the terms agnogenic myeloid metaplasia (AMM) or idiopathic myelofibrosis (IMF) lack accuracy since they are applied to both the prefibrotic hypercellular and advanced fibrotic stages of AMM or IMF. Consequently, Georgii replaced the terms...
prefibrotic AMM and IMF by chronic megakaryocytic granulocytic myeloproliferation (CMGM) and proposed in 1990 the Hannover Classification to distinguish 3 distinct primary CMPDs ET, PV and CMGM\(^\dagger\). The diagnosis of CMGM according to the Hannover classification\(^\dagger\) or prefibrotic CIMF-0 according to the Cologne criteria proposed by Thiele\(^\dagger\) (figure 3) is based on 3 main features.

First, the presence of large megakaryocytes with immature cytoplasm and immature cloud-like nuclei not seen in ET and PV. Second, increased granulopoiesis but never disturbed in maturation. Third, erythropoiesis is usually relatively decreased\(^\dagger\). The Hannover classification uses the term CMGM\(^\dagger\) and the Cologne classification uses the term prefibrotic chronic IMF (CIMF-0) for the third entity of prefibrotic MPD\(^\dagger\). The Hannover and Cologne classifications\(^\dagger\) are based on specific bone marrow histology (pathological) features for the three prefibrotic CMPDs true ET, PV and CIMF-0 at the bone marrow level, and have been taken over in 2001 by the WHO\(^\dagger\). Michiels and Thiele subsequently proposed the European clinical and pathological (ECP) criteria for the diagnosis of the Ph-negative MPDs to describe the full spectrum of clinical, laboratory, and WHO bone marrow features for true ET, early and overt PV and CIMF-0 (figure 3)\(^\dagger\). In a recent joint effort by clinicians and pathologists, we added the molecular marker and updated the European clinical, molecular and pathological (ECMP) criteria to extend PVSG criteria by including WHO bone marrow features and the use of laboratory and molecular markers for the diagnosis and staging of the three prefibrotic MPDs true ET, PV and CIMF (tables 2, 3 and 4)\(^\dagger\). These ECMP criteria allow a cross talk between clinicians and pathologists to translate PVSG clinical criteria (table 1) by including WHO bone marrow features and new biological and molecular markers (figures 2, 3, tables 2, 3 and 4) to reach agreement on diagnosis, classification and staging of the MPDs\(^\dagger\).

For the diagnosis of MPD bone marrow trephine biopsy specimens should be embedded in paraffin or plastic, both with its technical limitations. Paraffin requires decalcification with EDTA (preferable, allows reasonable DNA quality) or acid etch-solution\(^\dagger\). The specimens should have at least 4 evaluable bone marrow spaces with hematopoiesis. Recommended stains include: hematoxylin and eosin (H&E), Giemsa (3 μm sections), periodic acid-Schiff (PAS); Perls for estimation of hemosiderin content; chloro-acetate esterase (Leder) for identification of granulocytic differentiation; silver stain for reticulin; and trichrome-Masson for collagen staining. Immunostains of paraffin embedded specimens should include glycophorine C for erythropoiesis, myeloperoxidase for granulopoiesis, CD42b, CD61 or FVIII-related antigen for megakaryocytes; CD34 for CD34-positive blasts and CD117 for myeloid differentiation. Regarding MPD, clinicians want to receive from their pathologist\(^\dagger\) a detailed report according to the Cologne bone marrow evaluation form\(^\dagger\). According to WHO and ECMP criteria, increase and loose clustering of enlarged mature megakaryocytes with hyperlobulated nuclei in a normocellular bone marrow and platelet count >400 x10\(^9\)/l represent the hallmark of true ET (table 2, figures 2 and 3). In true ET there is no proliferation or immaturity of granulopoiesis or erythropoiesis. In congenital and acquired erythrocytosis and in reactive thrombocytosis the megakaryocytes are of normal size and morphology and there is no tendency to cluster. A typical histopathological ET picture of the bone marrow excludes RT and distinguishes true ET from early PV, CIMF-0, CIMF-1 (figure 2), and thrombocytosis associated with atypical MPD, MDS, refractory anemia with increased ringed sideroblasts (RARS-T) or Ph\(^+\)-positive thrombocytosis in CML. The characteristic increase and clustering of small and enlarged pleomorphic megakaryocytes and increased erythropoiesis with increased granulopoiesis and increased cellularity (80-100%) are the diagnostic characteristics of classic PV with increased hematocrit >0.51, low serum EPO and/or JAK2\(^V617F\) (tables 4 and 6) distinguishing it from congenital and secondary erythrocytosis. A typical histological PV picture with moderately increased bone marrow cellularity (according to age) is seen in patients with early PV mimicking ET or “forme fruste” PV featured by platelet count >400x10\(^9\)/l and hematocrit <0.51, low serum EPO and/or the presence of the JAK2\(^V617F\) mutation (tables 4 and 6). A typical histological PV bone marrow picture is also seen in the early erythrocythemic stage 1 PV featured by increased hematocrit >0.51, normal platelet count <400 x10\(^9\)/l, normal spleen, low serum EPO and/or the presence of the JAK2\(^V617F\) mutation (tables 4 and 6).

According to Hannover\(^\dagger\), Cologne\(^\dagger\), WHO\(^\dagger\) and ECMP criteria\(^\dagger\), CIMF-0 is characterized by hypercellularity of the bone marrow (60-100%) due to increased granulopoiesis, relative decrease of erythropoiesis and the presence of dense clusters of immature megakaryocytes with maturation defects.
of cytoplasm and nuclei with bulky nuclei showing clumpy lobuli and irregular roundish shaped forms (so-called cloud-like or clumsy nuclei), which are almost never seen in ET and PV (figures 2 and 3)\textsuperscript{57-73}. The degree of dysmegakaryopoiesis in CIMF-0 may range from slight maturation defect of megakaryocytes with no cloud-like nuclei to manifest maturation defects of megakaryocytes with typical cloud-like nuclei or a mixture of both. The risk of CIMF-0 to transform into early CIMF-1 and subsequent CIMF-2/3 with extramedullary hematopoiesis is clearly dependent on the degree of hypercellularity and on the degree of maturation defects of megakaryopoiesis\textsuperscript{70,71}. High quality histological bone marrow preparations in the hands of experienced hematopathologists are required to distinguish CIMF-0 from true ET and PV in about 85 to 90% of the cases, which is only feasible in centers of excellence (figures 2 and 3)\textsuperscript{57-73}. There are no studies that have examined the concordance between a number of pathologists who have used characteristic histological bone marrow features to assign cases to the prefibrotic stages of true ET, PV and CIMF-0 without knowledge of the clinical findings and biological MPD markers except age. Data on the very long-term natural history of patients with ECMP defined true ET, early PV and CIMF-0 as derived from large scale prospective studies are lacking. We don’t really know whether the early stages of PV patients are at no, low or high risk of progression to post-PV myelofibrosis or whether true ET never progress to CIMF as it is claimed. The bone marrow histology of CIMF-0 with slight dysmegakaryopoiesis can appear to overlap with early PV (hematocrit <0.51) presenting with a trilinear hypercellular bone marrow with relative increased granulopoiesis as compared to erythropoiesis and a similar degree of thrombocytopenia, leukocytosis, increased LAP-score and slight splenomegaly. The histology of CIMF-0 slight dysmorphic megakaryopoiesis may overlap with that of true ET in cases of very mild hyperplasia of granulopoiesis and/or a mixture of mild dysmorphic megakaryocytes and mature enlarged megakaryocytes with hyperploid nuclei. Diagnostic differentiation between true ET, early PV, CIMF-0 at the bone marrow level will be feasible in well equipped pathology laboratories by experienced trained pathologists in the majority of cases, but significant overlap (grey zones) between CIMF-0 with slight dysmegakaryopoiesis versus true ET or early PV due to a rather high inter-observer disagreement between hematopathologists in routine daily practice is very likely.

**Clinical relevance of ECMP criteria for diagnosis and staging of ET, PV and CIMF**

The PVSG-defined ET patients (table 1) comprise three ECMP-defined phenotypes of thrombocytopenia at the bone marrow level: true ET, early PV mimicking ET, and CIMF-0 or CIMF-1 without features of leuko-erythroblastosis in the peripheral blood\textsuperscript{25,70,71}. These three ECMP defined ET phenotypes do not differ significantly with regard to peripheral blood features, thrombocytopenia related clinical presentation and laboratory findings during long-term follow-up (table 7)\textsuperscript{70,71}. Therefore, patients with true ET and early PV and CIMF-0 mimicking ET are to be treated equally based on clinical risk stratification for thrombotic and bleeding complications irrespective of bone marrow features\textsuperscript{46}. The relevance of recognition of CIMF-0 and CIMF-1 lies in the increased risk of myelofibrotic transformation and decreased prognosis in terms of survival compared to true ET\textsuperscript{70,71}. The prognostic importance of the WHO/ECMP criteria is demonstrated in a large retrospective study of 476 PVSG-defined ET patients (platelet count >600 x10\textsuperscript{9}/l), who were reclassified according to the WHO bone marrow criteria: true ET in 167, CIMF-0 in 174 and CIMF-1 in 135\textsuperscript{71}. Mean age of true ET patients was 59 years, which is 8 to 10 years younger compared to CIMF-0 (67 years) and CIMF-1 (69 years) patients. The differences in relative 10 years survival rates: 99 ± 7.8% for true ET, 81 ± 11.7% for CIMF-0, and 67 ±17.8% for CIMF-1 patients, are significant due to an increased risk of myelofibrosis and splenomegaly during follow-up. In this retrospective “one-center-study” the majority of CIMF-0 patients have early stage disease without features of leuko-erythroblastosis (table 7), whereas CIMF-1 patients are a mixture of early and intermediated stage disease without and with leuko-erythroblastosis\textsuperscript{71} (as defined in tables 1 and 3 when applying ECMP criteria). The ECMP criteria for diagnosis and staging of CIMF patients do clearly separate early stage CIMF-0 and CIMF-1 without leuko-erythroblastosis from intermediate stage CIMF-1 and CIMF-2 with leuko-erythroblastosis of the peripheral blood (table 3)\textsuperscript{72,73}. Clinicians should realise that the overall survival curves of patients with early stage CIMF-0 or CIMF-1 and no leuko-erythroblastosis will be still as good as for newly diagnosed PV patients, and that the life expectancy of patients with CIMF-2 and CIMF-3 with leuko-erythroblastosis, splenomegaly and anemia will be significantly shortened (table 7).

A recent report of 116 PVSG-defined ET patients and reclassified based on WHO bone marrow cri-
aidia confirmed that such cohort of patients with the clinical picture of ET in fact comprises true ET in 19%, CIMF-0 in 21%, CIMF-1 in 37%, CIMF-2 in 12%, early PV in 8%, and unclassified MPD in 3%74. Median age of true ET and CIMF-0 patients was 54 and 52 years respectively, which is 7 to 14 years younger compared to CIMF-1 (59 years) and CIMF-2 (66 years), which points to the unexplored question whether true ET will progress to PV of CIMF after very long-term follow-up74. Thromboembolic events were equally frequent in ET, CIMF-0 and 1, but relevant life threatening events including acute myeloid leukemia, advanced CIMF and second malignancies were more frequent in CIMF-1 and 2 during long-term follow-up74.

In ECMP-defined true ET, progression into myelofibrosis grade 1 or 2 was not seen five years after diagnosis in two recent studies, but data on very long-term follow-up of more than 10 to 15 years are lacking75,76. Kvasnicka and Thiele calculated that the estimated risk of transformation within 3 years into clinically defined IMF (with anemia, leuko-erythroblastosis and extramedullary hematopoiesis) was 2.2% in PVSG-defined ET and 2.8% in WHO-defined CIMF-0 and -1 (nearly 1% per year)71. In a large series of 195 PVSG-defined ET patients and a median follow-up of 7.2 years, evolution into CIMF-2/3 (classic IMF with anemia, leuko-erythroblastosis and splenomegaly) occurred in 2.7% at 5 years, 8.3% at 10 years and 15% at 15 years (6.7% after a median of 8.3 years)77. In another retrospective study of 322 PVSG-defined ET, the cumulative risks of CIMF-2/3 and leukemia were 3.8% and 1.4% at 10 years, and 19.9% and 8.1% at 20 years respectively78. In these two studies of PVSG-defined ET patients, the overall survival was similar to that of the age-matched control population in the first decade, but significantly worse beyond the first decade of the disease. Applying ECMP criteria to PVSG-defined ET patients at time presentation will separate patients with true ET from early PV and CIMF-0 mimicking ET and therefore surely will become of prognostic importance to distinguish true ET from prefibrotic CIMF-0 and early fibrotic CIMF-1 in the context of new prospective clinical management studies.

Diagnostic work-up of patients with thrombocythemia in various MPD

Clinical features suspicious for thrombocythemia in various MPDs (true ET, early PV and CIMF-0) include a sustained increased platelet counts (>400 x10^9/l) in the absence of any cause for reactive thrombocytosis20,23,24,72,73. The presence of giant platelets in a peripheral blood smear is indicative for MPD and precludes reactive thrombocytosis. Sustained increase of platelet counts (>400 x10^9/l) associated with slight splenomegaly on echogram (>12cm), increased leukocytes (>12 x10^9/l) or LAP score with normal ESR is highly suspicious of myeloproliferative thrombocythemia. Clinical manifestations of thrombocythemia in various MPDs consist of microvascular circulation disturbances including atypical and typical TIAs, ocular ischemic attacks, erythromelalgia, splanchic or cerebral vein thrombosis41. Clinicians and pathologists should realise that PVSG-defined ET includes true ET, early PV mimicking ET, and thrombocythemia associated with CIMF-0 or CIMF-1 when ECMP criteria are applied (tables 1, 2, 3 and 4). The presence of numerous abnormal enlarged or giant mature megakaryocytes with hyperlobulated nuclei and preserved nuclear/cytoplasmic ratio or the presence of pleiomorphic small and enlarged or giant megakaryocytes with hypolobulated cloud-like nuclei, and/or the evidence of several clusters of enlarged megakaryocytes are the pathognonomic clues to the diagnosis of prefibrotic MPD (ET, PV or CIMF)3,19-33. The diagnostic work-up of patients with ET and thrombocythemia associated with CIMF according to ECMP criteria28,72,73 is based on positive criteria in peripheral blood and bone marrow (figure 5). These include:

1. Thrombocythemia patients should fulfil the peripheral blood (clinical) criteria for the diagnosis of thrombocythemia irrespective of bone marrow features (tables 2 and 3).

2. The screening for JAK2V617F as a first intention diagnostic test is very helpful in the diagnostic work-up of patients with suspected thrombocythemia in various MPDs, but only half of ET and CIMF patients carry this mutation.

3. Pretreatment bone marrow biopsy will allow clinicians and pathologists to diagnose the early stages of MPDs including JAK2V617F positive and JAK2 wild type thrombocythemia. The ECMP criteria classify the PVSG-defined ET (table 1) as: true ET (table 2); early PV mimicking ET (table 4); early stage CIMF-0 or CIMF-1 without features of leukoerythrocytosis and extramedullary hematopoiesis (table 3); and intermediate CIMF-1, 2 and 3 with features of leucoerythroblastosis79,80 (table 7).

4. The ECMP criteria distinguish thrombocythemia in various MPDs from thrombocythemia...
associated with Ph1-chromosome and bcr/abl positive chronic myeloid leukemia (CML) or myelodysplastic syndromes (MDS) including the so-called 5q-syndrome, which clearly differs from refractory anemia with ringed sideroblasts and significant thrombocytosis (RARS-T) (figure 5). Among 9 RARS-T patients in a recent study, 6 showed the presence of JAK2V617F mutation.

Comparing the laboratory features of JAK2V617F positive (in granulocytes) and JAK2 wild type PVSG-defined ET patients in the PT-1 study showed that JAK2V617F positive ET is characterized by higher values for hemoglobin, hematocrit, neutrophil counts, LAP score, by lower values for serum EPO levels, serum ferritin and MCV, and by increased cellularity of the bone marrow in biopsy material. This observation confirms the ECMP concept that JAK2V617F and EEC positive ET patients represent an early PV mimicking ET (“forme fruste” PV, stage 1 PV, table 5). As compared to JAK2V617F positive ET (early PV), JAK2 wild type ET patients had significantly higher platelet counts, normal serum EPO levels, a typical bone marrow picture of true ET, no features of early PV, and are at lower risk for the development of thrombotic complications. These data are in line with the hypothesis that JAK2V617F positive and JAK2 wild type ET patients at diagnosis represent two distinct entities with a related pathophysiology in the JAK-2/STAT signalling pathway but different molecular etiology similar to the gain of function mutation in the TPO or MPL genes causing hereditary ET.

**Diagnostic work-up of patients with polycythemia vera**

Suspected polycythemia (PV) with characteristic PV features include increased hematocrit (>0.51), increased erythrocytes (>6 x10^12/l), slight splenomegaly, increased leukocytes (>12 x10^9/l) or LAP score with normal ESR, increased platelets (>400 x10^9/l). PV patients usually show the presence of large platelet in peripheral blood smear. PV patients frequently present with headache, TIA’s, erythromelalgia, splanchic or cerebral vein thrombosis and microcytosis of erythrocytes due to iron deficiency. Patients with congenital or acquired erythrocytosis lack the clinical and laboratory features of MPD are usually asymptomatic. The presence of JAK2V617F mutation in granulocytes has a sensitivity of about 95% and positive predictive value of 100% for the diagnosis of PV in the context of absolute erythrocytosis (hematocrit >0.51 in males and >0.48 in females) and excludes congenital and secondary erythrocytosis (figure 6). Subsequent red cell mass measurement will distinguish apparent from absolute erythrocytosis but does not differentiate between PV and congenital or secondary erythrocytosis. In contrast, bone marrow histology not only differentiates trilinear hypercellularity in PV (figures 2 and 3) from isolated increase of erythropoiesis (figure 3) in congenital polycythemia and secondary erythrocytosis, but also significantly contributes to phenotyping and staging of PV patients.

The detection of JAK2V617F in granulocytes with sensitive PCR techniques as to play a key-role as a first intention diagnostic test for erythrocytosis (hematocrit >0.51), because it simplifies the diagnostic work-up of PV (table 4, figure 6). The presence of the JAK2V617F mutation combined with increased hematocrit (>0.51), and EEC or low serum EPO is diagnostic for PV without the need of red cell mass measurement (table 4), but is not enough to define the broad spectrum of PV phenotypes according to EMCP criteria (table 6). Since EEC is time consuming and difficult to establish in many (non-specialized) laboratories, clinicians and pathologists tend to replace it by the wide spread available bone marrow histology assessment as a gold standard criterion for the diagnosis of PV and its differentiation from the EMCP-defined true ET and prefibrotic or early fibrotic CIMF (tables 3 and 4, figures 2 and 3).

Comparing 45 JAK2V617F heterozygous and 13 homozygous PV patients showed that homozygote JAK2V617F PV patients displayed significantly higher hemoglobin at time of diagnosis, increased incidence of pruritus, higher PRV-1 transcripts in their blood granulocytes, and higher rate of fibrotic transformation. These observations indicate that pretreatment and follow-up bone marrow histology examinations will be helpful for proper staging of PV patients for reasons of prognosis assessment and therapeutic implications (table 6). In the context of a prospective clinical study to discriminate between early and advanced PV and to monitor disease progression to post-PV myelofibrosis, bone marrow histology, cytogenetic analysis and JAK2V617F mutation detection in granulocytes, and EEC should always be performed.

**Grading of myelofibrosis in myeloproliferative disorders (MPD)**

Myelofibrosis (MF) itself is not a disease because reticulin and collagen fibrosis are produced by pol-
yclonal fibroblasts as the consequence of cytokines released from the clonal granulocytic and megakaryocytic proliferative cells in both PV and CIMP\(^{19,97}\). The Baumeister scoring system of MF was developed on aspirated bone marrow samples, but proved to be not reliable for the proper grading of myelofibrosis in bone marrow biopsies by pathologist (table 8)\(^{98}\). The Manoharan system used silver stain according to Gordon and Sweet and scored the degree of reticulin in bone marrow biopsy in a completely different (table 8)\(^{99}\). A scoring system based on morphometric analysis (point intersection with an ocular grid) and quality of fibers (reticulin and collagen fibers) and the bone marrow fiber density (fine or course reticulin and some or course bundles of collagen) have been proposed by Georgii et al\(^{19,63,64}\) and Thiele et al\(^{65-68}\). All these different scoring systems for MF use different criteria for grading of reticulin and collagen, are subjective and not comparable by lack of strict criteria. A panel of experienced European pathologists and a USA expert reached a consensus on how to grade bone fibrosis in bone marrow biopsies of patients with CIMP or PV (EC, tables 3 and 8)\(^{100}\). Grading of MF was simplified by using four easily reproducible categories that included differentiation between reticulin and collagen\(^{100}\). According to defined standardized semiquantitative grading of reticulin and collagen fibrosis in the bone marrow, MF can reliably be graded at the pathological bone marrow level as 0 in prefibrotic, as 1 in early fibrotic, as 2 in classical fibrotic and as 3 in classical sclerotic CIMP (table 3 and 8)\(^{100}\). Myelofibrosis is rare in ET and does occur in about one third of PV and in the majority of CIMP-0 patients during long-term follow-up\(^{63,64,69,70,76}\).

Conclusion: The diagnosis CIMP according to WHO bone marrow features does not distinguish between CIMP-0/CIMP-1 without leuko-erythroblastosis versus CIMP-1/CIMP-2 with leuko-erythroblastosis when ECMP criteria are applied (tables 3 and 7) indicating the need to use clinical score assessment on top of bone marrow features for prognosis prediction.

References


The Targets of Therapy in Polycythemia Vera and Thrombocythemia

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The most common myeloproliferative disorders (MPDs) include Polycythemia Vera (PV), Essential Thrombocythemia (ET) and Idiopathic Myelofibrosis (IMF). These are clonal diseases resulting from a transformation of a multipotent hematopoietic stem cell that leads to overactive hemopoiesis and bone marrow fibrosis. The clinical course of PV and ET is marked by significant thrombotic complications and a variable risk to evolve into acute myeloid leukemia. Polycythemia vera (PV) and Essential Thrombocythemia (ET) are Chronic myeloproliferative disorders (MPDs) in which morbidity and mortality are more frequently due to thrombohemorrhagic diathesis and a variable incidence of progression to myelofibrosis (MF) or acute leukemia.

Randomized clinical trials performed in USA and Europe have shown that cytoreductive treatment of blood hyperviscosity, chemotherapy and low-dose aspirin have dramatically reduced the number of thrombo-hemorrhagic episodes and substantially improved survival. However, there is a concern that certain myelosuppressive drugs accelerate the disease progression to acute leukemia. Thus, to minimize this risk, the best strategy in PV and ET is to limit the cytotoxic drugs to patients stratified on the basis of their risk for developing vascular events. The important risk factors are an age of 60 years or more and previous vascular events whereas hemorrhagic complications are paradoxically associated with extreme thrombocytosis.

As compared to PV and ET, IMF has the worst prognosis with a median survival of 3-5 years. A prognostic score system was developed where the presence of leukocytosis, leukopenia or anemia was used to identify three groups of patients with different survival, from 1 to 8 years. Conventional therapy in this disease is palliative and includes many drugs in addition to supportive therapy to improve anemia, thrombocytopenia and progressive splenomegaly. Recently, an experimental approach with hematopoietic stem cell transplantation after a reduced intensity conditioning regimen is offered in the younger patients and the results seem promising.

A new avenue for the treatment of MPDs was opened by the recent identification of an acquired mutation of the JAK2 gene (V617F) in 90% of patients with PV and in about half the patients with ET and IMF. The consequence of this mutation is a constitutive tyrosin kinase activity of JAK2 resulting in proliferative and survival advantage of hematopoietic progenitor cells. This finding is being explored in terms of diagnosis, prognosis and therapy. The mutational status and the allele burden in granulocytes have been correlated with survival, specific disease symptoms and the evolution towards leukemia. The results on its prognostic value are so far inconclusive and do not provide new stratification for therapy. Novel JAK2 inhibitors are being developed and promising results have been obtained with some of these compounds. In this context, it should be mentioned that JAK2V617F is not an essential pathogenetic component for MPDs other than PV and might not be the initial clonogenic event. There are data to admit that other mutations may cooperate to determine the phenotype. Thus, these drugs should be tested in all MPDs and not limited to those with JAK2 mutation. The inclusion criteria for phase II studies should consider high risk patients with poor survival and those with myelofibrosis (primary or secondary to PV or ET) should be the first candidates for testing these experimental drugs.
CML: Case Closed?

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Chronic myeloid leukaemia (CML) is a malignant haemopoietic stem cell disorder characterised by the t(9;22)(q34;q11) reciprocal chromosomal translocation, the functional consequence of which is the Bcr-Abl oncoprotein. In 1998, imatinib, a tyrosine kinase inhibitor belonging to the 2-phenylaminopyrimidine group of pharmacological compounds, was introduced into the armamentarium of drugs for the treatment of CML, and has since revolutionised its management. Imatinib has a high affinity for the ATP-binding site of Abl, in addition to other kinases such as PDGFR, Kit and Arg, and clinical trials have validated the promise of this molecular targeted therapy. In the more advanced phases of CML, imatinib was able to induce a major (complete or partial) cytogenetic response in 16-60% of patients. In a phase III trial comparing imatinib with interferon-α plus cytosine arabinoside in newly diagnosed chronic phase CML, 85% of patients treated with imatinib attained a major cytogenetic response (MCyR) after a median follow-up of 19 months, compared to 22% in patients treated with the latter combination. A recent update has shown a further increase in MCyR of up to 92% of patients in the imatinib arm after a median follow-up of 54 months. In view of its high efficacy and low toxicity, imatinib has now replaced interferon-α as frontline treatment for CML patients who are not eligible for allogeneic stem cell transplantation.

Clinical resistance to imatinib

The efficacy of imatinib in CML is remarkable, but the development of resistance and the persistence of minimal residual disease have dampened the initial enthusiasm for this much heralded ‘magic bullet’. Resistance can be defined on the basis of its time of onset. Primary resistance is a failure to achieve a significant haematological or cytogenetic response, whereas secondary or acquired resistance is the progressive reappearance of the leukaemic clone after an initial response to the drug. Resistance is also defined on the basis of clinical and laboratory criteria used for detection of leukaemia, which includes haematological, cytogenetic and molecular resistance. Haematological resistance is a lack of normalisation of peripheral blood counts and spleen size; cytogenetic resistance is a failure to achieve a MCyR, i.e. less than 35% Philadelphia (Ph) chromosome positivity; and molecular resistance represents the failure to achieve or the loss of complete or major molecular response (MMR). MMR can be defined as a 3 or more log-reduction of $\text{BCR-ABL} / \text{control gene}$ ratio from a laboratory standardised baseline or an international scale converted $\text{BCR-ABL} / \text{control}$ gene ratio of < 0.1%. The attainment of a MCyR or MMR has an impact on progression-free survival. Early chronic phase patients who achieved a MCyR after 12 months of imatinib had a 96% rate of survival without progression to accelerated phase or blast crisis at 54 months, compared to 81% who did not achieve a MCyR. Achieving a complete cytogenetic response (CCyR) and a MMR in this cohort of patients translated to a 100% rate of survival without progression to the more advanced phases, compared to 95% in those who achieved a CCyR but not a MMR, and 89% in those who do not achieve a CCyR. Recently, the European Leukaemia Net refined response criteria allowing...
resistance to be categorised into 2 groups, ‘sub-optimal response’ and ‘failure to respond’ (Table I). Continuing imatinib treatment is unlikely to be beneficial in those with failure to respond, while the suboptimal responders may still have a benefit in continuing, although with a less favourable long term outcome 8.

Molecular basis of resistance

Ever since the first reports of resistance were described in 2000, the mechanisms of resistance to imatinib have been extensively studied and three major mechanisms have been identified. The two most common affect the BCR-ABL gene itself, namely mutations in its tyrosine kinase domain and overexpression of the Bcr-Abl protein due to amplification of the BCR-ABL gene 9,10,11,12,13,14,15,16,17,18,19,20,21,22,23,24,25,26. The third mechanism is less well characterised and understood, and is represented by phenomena which lead to resistance independent of Bcr-Abl. These include upregulation of the drug efflux pumps 27,9,28,29,30, downregulation of drug influx transporters 31,32,33, binding of the α1-acid glycoprotein (AGP) 34, overexpression of Lyn, a Src kinase 35, and other Bcr-Abl-independent mechanisms 36.

Mutations in the Abl kinase domain

The emergence and selection of clones exhibiting point mutations in the Abl kinase domain is the most frequently identified mechanism of resistance in patients treated with imatinib and is more common in acquired than in primary resistance 37. These mutations are not induced by imatinib, but rather, just like antibiotic resistance in bacteria, arise through a process whereby the drug itself selects for rare pre-existing mutant clones, which gradually outgrow drug-sensitive cells 38.

Mutations can be categorised into 4 groups: (i) those which directly impair imatinib binding; (ii) those within the ATP binding site; (iii) those within the activation loop, preventing the kinase from achieving the conformation required for imatinib binding; and (iv) those within the catalytic domain.

The substitution of the amino acid threonine with isoleucine at position 315 of the Abl protein was the first mutation to be detected in resistant patients 10. Based on the crystal structure of the catalytic domain of Abl complexed to a variant of imatinib 39, this substitution was predicted to reduce the affinity for the drug in two ways. Firstly, the oxygen atom provided by the side chain of threonine 315 is not present, and this prevents the formation of a hydrogen bond with the secondary amino group of imatinib. Secondly, isoleucine contains an extra hydrocarbon group on its side chain and this sterically inhibits the binding of the inhibitor 10. Another amino acid that makes contact with imatinib is phenylalanine 317, and its mutation to leucine also leads to resistance.

Mutations can also cluster within the ATP-binding loop (phosphate or P-loop). This domain is a highly conserved glycine-rich sequence that spans amino acids 248-256 and interacts with imatinib through hydrogen and van der Waals bonds 37. These mutations modify the flexibility of the P-loop and destabilise the conformation required for imatinib binding 14. Apart from imatinib insensitivity, a feature of clinical relevance is that imatinib-treated patients who harbour P-loop mutations have been suggested to have a worse prognosis than those with non-P-loop mutations 19. However, this has not yet been confirmed in larger series.

The activation loop of the Abl kinase begins at amino acid 381 with a highly conserved motif of 3 amino acid residues (Aspartate-Phenylalanine-Glycine). This region of the kinase can adopt a closed (inactive) or an open (active) conformation. Imatinib forces Abl into the inactive conformation and is incapable of binding to the active configuration 40. Mutations in the activation loop may disturb the energetic balance required to stabilise the closed conformation of the loop and thus favour the open, active conformation 14.

Finally, some amino acid substitutions cluster in the catalytic domain, a region that has a close topologic relation to the base of the activation loop. Mutations in this region can also influence the binding of imatinib 14.

At least 73 different point mutations leading to substitution of 50 amino acids in the Abl kinase domain have been isolated from CML patients resistant to imatinib so far, and this number is likely to increase with more sensitive methods of detection.

The detection of a Ph-positive clone harbouring an Abl kinase domain mutation is associated with resistance to imatinib and may be associated with progression to a more advanced phase disease 19,26. Using highly sensitive assays, mutations have also been detected in patients in complete cytogenetic response, and in imatinib-naïve advanced phase...
but not chronic phase patients 24,25. However, detection of mutations in these patients did not always result in progressive disease while on imatinib. It is likely that mutant clones in the presence of low leukaemic burden or low levels of mutant clones do not have a similar clinical impact as clones which are detected when disease burden is rising or high 25,41. Furthermore, the probability of detecting a clone is low when BCR-ABL transcript levels are stable or declining 21.

**Bcr-Abl overexpression**

Overexpression of the Bcr-Abl protein due to amplification of the BCR-ABL gene was first observed in vitro when resistant CML cell lines were generated by exposure to gradually increasing doses of imatinib 27,9,42,43. This phenomenon has been reported in a relatively small proportion of patients, with an overall percentage of 18% 17,10,37, but this may be an underestimate if its detection is only based on the cytogenetic findings of Ph chromosome duplication. In one study, 3 out of 11 CML patients in blast crisis who relapsed after initially responding to imatinib were shown to have multiple copies of the BCR-ABL gene by fluorescence in situ hybridisation (FISH) 10. In another study, 7 out of 55 patients showed a more than 10-fold increase in BCR-ABL transcript levels and 2 out of the 32 patients evaluated were found to have genomic amplification of BCR-ABL by FISH 37. In the latter 2 patients, resistance was primary and not acquired. Overexpression of Bcr-Abl leads to resistance by increasing the amount of target protein needed to be inhibited by the therapeutic dose of the drug. It is also possible that a transient overexpression of Bcr-Abl may be an early phenomenon in the establishment of imatinib resistance, preceding the emergence of a dominant clone with a mutant kinase domain, as suggested by kinetic studies in cell lines 44.

**Drug efflux and influx transporters**

Multidrug resistance (MDR) due to cross-resistance of mammalian cells to a number of anticancer agents following exposure to one such drug is a well described mechanism of resistance in cancer therapy. This is mediated by an increased expression at the cell surface of the MDR1 gene product, Pgp, an energy dependent efflux pump, which reduces intracellular drug concentrations and leads to ineffective levels of the drug reaching its target 45,46. Imatinib and other tyrosine kinase inhibitors are substrates of Pgp, and the intracellular levels of imatinib were shown to be significantly lower in Pgp-expressing cells 47,48,31,49,50. An imatinib-resistant CML cell line generated by gradual exposure to increasing doses of the drug was shown to exhibit Pgp overexpression, and MDR1 overexpression in CML cell lines also confers resistance to imatinib 9,28. Pgp overexpression has not been reported in patients who are resistant to imatinib. However, the addition of a Pgp pump inhibitor, PSC833, to cultures of imatinib-treated cells from drug-resistant CML patients produced a significant decrease in colony formation, thus suggesting that MDR1 overexpression may play a role in clinical imatinib resistance 28.

Recently, two other drug transporters, breast cancer resistance protein (BCRP)/ABCG2 and human organic cation transporter 1 (hOCT1), have been implicated as possible mechanisms for promoting imatinib resistance. Imatinib has been variably reported to be a substrate and/or an inhibitor for the BCRP/ABCG2 drug efflux pump which is overexpressed in many human tumours and also found to be functionally expressed in CML stem cells 51,52,53,54,29,55,30. The drug transporter, hOCT1 mediates the active transport of imatinib into cells, and inhibition of hOCT1 decreases the intracellular concentration of imatinib, which may predict for a less favourable molecular response 31,33. The hOCT1 gene was also found to be expressed in significantly higher levels in patients who achieved a complete cytogenetic response to imatinib than in those who were more than 65% Ph chromosome positive after 10 months of treatment 32. This would suggest that patients with low baseline expression of hOCT1 may not achieve a complete cytogenetic response because of insufficient intracellular levels of imatinib.

**Bcr-Abl independent mechanisms**

The Src family kinases, Lyn and Hck, are activated in BCR-ABL-expressing cell lines. Lyn is overexpressed and activated in an imatinib-resistant CML cell line generated by incubation of the parental line in increasing concentrations of imatinib and in samples from CML patients who were resistant to imatinib 35. Lyn suppression by a Src kinase inhibitor resulted in reduced proliferation and survival of the imatinib-resistant but not the sensitive cell line 35.

Microarray analysis have shown that transcripts from genes with anti-apoptotic or malignant transformation properties and with involvement in
signal transduction/ transcriptional regulation are overexpressed in CML cells innately resistant to imatinib. This would suggest that pathways downstream of Bcr-Abl and independent of its kinase activity may be important factors which confer resistance to imatinib 36. The phosphatidylinositol-3 (PI-3) kinase/Akt pathway is an important downstream pathway activated by Bcr-Abl and is essential for the proliferation of BCR-ABL-positive cells 56. The mammalian target of rapamycin (mTOR) is a serine-threonine kinase activated by the PI-3 kinase. Treatment with imatinib was shown to activate the PI-3 kinase/Akt/mTOR pathway and this activation was important in mediating cell survival during the early development of imatinib resistance before overt resistance developed 57.

**Overcoming imatinib resistance**

Since the discovery of imatinib resistance it became clear that there is an urgency to discover and develop novel compounds or combinations capable of circumventing it. A number of potent Abl kinase inhibitors with in vitro and in vivo activity in wild-type and Abl kinase mutant Bcr-Abl cell lines have been identified. Two of these compounds are now in clinical trials. Pre-clinical studies have also provided evidence that combination therapy may have an important role in preventing or combating imatinib-resistance.

**Dual Src/Abl kinase inhibitors**

The disruption of the proto-oncogene Src is associated with the pathogenesis of human cancers 58. Synthetic small molecule inhibitors of Src-family kinases have been developed and these compounds, eg, PD180970, AP23464, SKI606, CGP76030, BMS-354825, also inhibit Bcr-Abl, Kit and PDGFβ receptors, and have in vitro anti-proliferative activity in imatinib-sensitive and -resistant CML cells 59,60,61,62,63,64. Dasatinib (BMS-354825, Bristol Myers Squibbs) was found to be more potent than imatinib, and was capable of inhibiting the proliferation and kinase activity of wild type Bcr-Abl cell lines at picomolar concentrations 65. Similar to previously developed Src/Abl kinase inhibitors, dasatinib is also active against the imatinib-resistant active conformation of the kinase domain. In vitro assays revealed that dasatinib inhibited the kinase activity and proliferation of 14 out of 15 clinically relevant Bcr-Abl mutant cell lines. However, the T315I mutant remained resistant, even at micromolar concentrations of the drug. In vivo studies in murine models have confirmed the activity of dasatinib in inhibiting the leukaemic cell growth and prolonged the survival of mice harbouring the wild type Bcr-Abl and the M351T, but not the T315I mutant 63.

Clinical trials of dasatinib in imatinib-resistant and -intolerant CML and Ph chromosome-positive acute lymphoid leukaemia (Ph+ALL) are currently in progress, and the haematological and cytogenetic responses are summarised in Table II 66,67,68,69,70. Dasatinib is well tolerated but grade 3-4 myelosuppression is common, especially in the advance phases. Non-haematological side effects include diarrhoea, nausea, headache, peripheral oedema and pleural effusion.

**Second generation Abl kinase inhibitors**

The N-methylpiperazine moiety was originally incorporated into imatinib to improve its solubility and oral bioavailability. Substitution of this amide moiety with alternative binding groups, while maintaining H-bond interactions to Glu286 and Asp381, led to the discovery of a more potent compound, nilotinib (AMN107, Novartis). Nilotinib also inhibits the activity of Arg, Kit, and PDGFα and β receptors, but not Src kinase. At submicromolar concentrations it is 10 to 50 times more potent than imatinib in inhibiting the proliferation and autophosphorylation of wild-type Bcr-Abl cell lines and of most of the clinically relevant Bcr-Abl mutants, except the T315I mutant. Nilotinib was also superior to imatinib in reducing leukemic burden and prolonging the survival of mice transplanted with marrow transduced with wild-type Bcr-Abl, the M351T and E255V mutants 71. Results from phase I clinical trials with nilotinib are summarised in Table III 72.

**Substrate-competitive inhibitors**

Adaphostin is a tyrphostin which alters the binding of peptide substrates rather than the ATP-binding site. Imatinib-resistant cell lines were shown to remain sensitive to the inhibitory effects of adaphostin 73. Adaphostin does not target Bcr-Abl specifically but it selectively inhibited colony formation from primary CML cells 73. It has also been shown to induce cytotoxicity by generation of reactive oxygen species 74.

The resistance of the T315I mutant to the Src/Abl kinase inhibitors and nilotinib poses a therapeutic challenge, and it is likely that this mutant will remain insensitive to other ATP-competitive inhibitors. A substrate-competitive inhibitor of
Bcr-Abl, ON012380, was recently reported to have potent in vitro inhibitory activity in cell lines expressing wild-type Bcr-Abl and all the Bcr-Abl mutants, including the T315I mutant. The activity against the T315I mutant was confirmed in vivo in mice expressing this form of Bcr-Abl protein where treatment with ON012380 caused a decrease in leukaemic cells 75.

**Allosteric inhibitors**

A recent class of Bcr-Abl inhibitor compounds was uncovered by differential cytotoxicity screen in a 384-well format of approximately 50,000 combinatorially derived kinase-directed heterocycles 76. This is a class of compounds which exert their activity through a newly described allosteric, non-ATP competitive mechanism, potentially involving binding to the myristate pocket in the C-loop of the Bcr-Abl kinase domain.

**Other compounds with activity against imatinib-resistance**

Many compounds have now been described to exhibit in vitro activity against imatinib-resistant cells. These include the inhibitors of the Bcr-Abl chaperone heat shock protein 90, geldanamycin and 17-allylamino-geldanamycin (17-AAG)77; the MEK kinase inhibitor, PD18435272; the cyclin-dependent kinase inhibitor, flavopiridol75; the histone deacetylase inhibitors, LBH58980, suberoylanilide hydroxamic acid (SAHA) and sodium butyrate81; the proteasome inhibitor, bortezomib82; and the dual Abl/Lyn kinase inhibitor, NS-18783.

Other small molecule compounds have also been identified to have in vitro anti-proliferative activity against the T315I mutant and these include a phosphoinositide-dependent kinase-1 inhibitor, OSU-0301284; an Aurora kinase inhibitor, VX-68085; a p38 inhibitor, BIRB-79685; and an Abl kinase inhibitor, SGX-7043086.

**Combination therapy with inhibitors of effectors in Bcr-Abl downstream pathways**

Targeting Bcr-Abl downstream pathways is another attractive strategy for overcoming and possibly preventing resistance. Farnesyl transferase inhibitors (FTI) inhibit protein farnesylation and antagonise the oncogenicity of Ras, a protein that plays a central role in leukaemogenic transformation by Bcr-Abl. Lonafarnib (SCH66336, Schering-Plough) is an FTI which inhibits the proliferation of Bcr-Abl wild type, overexpressing and T315I mutant cell lines. While the drug itself did not induce apoptosis, it enhanced imatinib-induced apoptosis in the first two cell lines but not the T315I mutant 87. Lonafarnib also enhances imatinib-induced cytotoxicity in primitive quiescent CML cells, a population of cells known to persist in vitro in imatinib-treated primary CML cells 88. Nitrogen-containing bisphosphonates (e.g., zoledronic acid), via their inhibition of Ras prenylation, have also been effective in inducing apoptosis and inhibiting proliferation of imatinib-sensitive and -resistant CML cells 89,90.

The PI-3 kinase/Akt/mTOR pathway effectors are also candidates for targeted molecular therapy. The combination of the mTOR inhibitor, rapamycin or its derivative, RAD001, with imatinib was effective in overcoming imatinib-resistance in cell lines which overexpressed Bcr-Abl or harboured mutants which retained a moderate sensitivity to imatinib. However this combination was not effective in mutants which were highly resistant to imatinib, for example, T315I and E255K 91,92. A novel phosphoinositide-dependent kinase-1 inhibitor, OSU-03012, acting via an Akt-dependent mechanism, was shown to synergise with imatinib in inducing apoptosis and inhibiting proliferation in both the T315I and E255K mutants 84.

The Jak-STAT pathway is the third major pathway downstream of Bcr-Abl. Mycophenolic acid (MPA), an inosine monophosphate dehydrogenase inhibitor that depletes intracellular guanine nucleotides, reduced phosphorylation of STAT5 and S6 ribosomal protein, a substrate downstream of mTOR. When combined with imatinib, MPA produced synergistic antiproliferative and pro-apoptotic effects in imatinib-sensitive CML cells. However the effect on imatinib-resistant cells was not determined 93.

**Conclusion**

The introduction of imatinib has represented a major achievement for the treatment of CML but the development of resistance to this drug has become a potential therapeutic dilemma. It is clear, therefore, that monotherapy with imatinib may not be the best option in CML, or at least not for all patients. Therapeutic approaches to circumvent the problem of imatinib-resistance are now possible with novel compounds and combinations being investigated pre-clinically and clinically. Dasatinib and nilotinib represent the first of the newer generation tyrosine kinase inhibitors to have a good safety profile and efficacy in imatinib-resistant
and -intolerant CML patients. It is not possible, currently, to determine which is more effective or less toxic. However, subclones with hitherto unseen Bcr-Abl mutants will probably develop in response to these new small-molecule inhibitors, leading again to resistance to these compounds. One strategy to delay or suppress the emergence of these mutants is, therefore, to inhibit Bcr-Abl downstream signalling pathways and, possibly, to achieve a synergistic combination with the newer Abl kinase inhibitors.

References


The Molecular Pathogenesis of MDS

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Abstract
Clonal disorders of hematopoiesis, such as myelodysplastic syndromes (MDS) and myeloproliferative diseases (MPD), affect both hematopoietic stem cells and progenitor cells within the erythroid, platelet and granulocytic lineages and can have devastating consequences in children and adults. The genetic features of these diseases often include clonal, nonrandom chromosomal deletions (e.g., 7q-, 5q-, 20q-, 6q-, 11q- and 13q-) that appear to inactivate tumor suppressor genes required for the normal development of myeloid cells (reviewed in Bench1 and Fenaux2). These putative tumor suppressors have proved to be much more difficult to identify than oncogenes activated by chromosomal translocations, the other major class of chromosomal lesions in MDS and MPD.3 Although MDS and MPD are almost certainly caused by mutations in stem/progenitor cells,4 the role of inactivated tumor suppressor genes in this process remains poorly understood. In a small portion of myeloid diseases, mutations have been identified in genes encoding factors known to be required for normal hematopoiesis, such as PU.1, RUNX1 and c/EBPα, and implicating these genes as tumor suppressors.5-7 In addition, we have recently shown that CTNNB1 (α-catenin) is deleted on one allele and epigenetically silenced on the other allele in MDS patients with (del)5q.7,14 Nonetheless, the identities of most deletion-associated tumor suppressors in these diseases remains elusive, despite complete sequencing of the human genome. The deleted regions detected by cytogenetic methods are generally very large, containing many hundreds of genes, thus making it hard to locate the critical affected gene or genes. It is also unclear whether dysfunctional myelopoiesis results from haploinsufficiency, associated with the deletion of one allele, or from homozygous inactivation due to additional point mutations or microdeletions of the retained wild-type allele. In general MDS have proved surprisingly resistant to conventional treatments. Targeted therapeutic advances in MDS will likely depend on a full comprehension of underlying molecular mechanisms, in particular the tumor suppressor genes lost through clonal, nonrandom chromosomal deletions, such as the 7q- and (del)5q.

Introduction
Myelodysplastic syndrome (MDS) refers to a group of clonal disorders characterized by trilineage defects in hematopoiesis, including the erythroid, granulocytic, and megakaryocytic lineages. Although clonal, it is sometimes considered a premalignant condition that often progresses to acute myeloid leukemia (AML), when additional genetic abnormalities are acquired.8-10 Overall, MDS affects approximately 1 in 500 persons over 60 years of age, making it the most common hematologic malignancy in this age group;11 it may develop at any age, including childhood. As a complication associated with aggressive treatment of other cancers, MDS shows a high correlation with exposure to radiation, alkylating agents or topoisomerase II inhibitors.12-17 MDS often develops following autologous bone marrow transplantation, affecting 20% of patients with non-Hodgkin lymphomas who received bone marrow transplants.13,15,18,19 The prognosis for patients with primary or secondary MDS remains poor, especially in the elderly. MDS usually requires allogeneic bone marrow transplantation for permanent cure, but unfortunately, older patients cannot generally tolerate this procedure, leaving them without effective alternatives. In a study of adult patients with primary MDS, only 6% were alive and in remission 7 years after diagnosis.20 Children with MDS who undergo bone marrow transplantation have a 58% survival rate after 3 years, as compared with an average survival of only 0.9 years for those who do not receive a transplant.21 These bleak statistics underscore the importance of understanding MDS at the molecular level in order to expand the repertoire of biologically based therapies. Improvement in the treatment of acute promyelocytic leukemia (M3 subtype), using all-trans-retinoic acid to induce differentiation of malignant promyeloblasts, illustrates the potential of such strategies.22-24
Although MDS has been recognized as an important disease for more than 50 years, its molecular pathogenesis and the molecular basis for its progression to AML remain largely undefined. A model of MDS molecular pathogenesis has been proposed whereby a normal hematopoietic stem cell acquires successive genetic abnormalities that ultimately lead to malignant transformation and clonal expansion. Evidence for clonality in MDS comes primarily from nonrandom X-inactivation studies performed on the bone marrow cells of female patients with MDS. These studies demonstrate clonal involvement of hematopoietic cells in this disorder. Early mutations in stem cells may cause differentiation arrest leading to dysplasia, whereas subsequent defects affecting myeloid cell proliferation may cause the clonal expansion of aberrant cells and frank AML. Although many chromosomal abnormalities have been detected in MDS (e.g., 20q- and monosomy 7), the genes involved are yet to be identified, and it is unknown whether these genetic aberrations are initial events leading to the development of MDS or are secondary events.

Myeloproliferative disorders (MPD), such as chronic myeloid leukemia (CML) and myeloid metaplasia with myelofibrosis (MMM), are hematopoietic stem cell diseases characterized by uncontrolled growth of granulocytes and other hematopoietic cells, resulting in clonal expansion of those lineages. A role for mutationally activated tyrosine kinase genes has now been established in most cases with MPD (Fig. 1). Recently, following earlier discoveries of mutant tyrosine kinases in CML, chronic myelomonocytic leukemia, hypereosinophilic syndrome and systemic mast cell disease, four groups reported a specific activating mutation in the tyrosine kinase JAK2 in three distinct forms of MPD — polycythemia rubra vera (PRV), essential thrombocytemia (ET), and MMM.4,25-27 Despite these advances in the identification of tyrosine kinase oncogenes, it has been difficult to identify the tumor suppressors whose inactivation contributes to the pathogenesis of MPD and the chromosomal regions of deletion are often shared between MDS and MPD (Fig. 1). Patients with neurofibromatosis type I, with inactivation of the neurofibromin 1 (NF1) tumor-suppressor gene, frequently develop juvenile myelomonocytic leukemia (JMML); however, the onset of this and other types of MPD is commonly associated with complex karyotypes indicating the involvement of additional genetic pathways in the pathogenesis of MPD.28 Clearly, a major advance in the fields of MDS and MPD will be made by the identification of additional mutations involved in the development and progression of these families of diseases.

**Chromosomal Loss and Malignant Progression in Human Myeloid Diseases**

**Figure 1.** Common chromosomal deleted regions, duplications and other abnormalities associated with (A) MDS, (B) MPD and (C) activated kinases associated with MPD. The pie charts represent the percentages of patients with the indicated cytogenetic abnormalities estimated from the following studies: **Fig. 1A.** Based on data in List AF et al, 2004, and **Fig. 1B** from based on data in Bench1. 

Abbreviations for **Fig. 1C:** CML, chronic myelogenous leukemia; CMML, chronic myelomonocytic leukemia; EMS, 8p11 myeloproliferative syndrome; CEL, chronic eosinophilic leukemia; SM, systemic mastocytosis.; PV, polycythemia vera; ET, essential thrombocytemia; IMF, idiopathic myelofibrosis. (Reproduced with permission from a slide provided by Dr. Anthony Green).
mal karyotypes, (ii) balanced chromosomal aberrations leading to the generation of fusion oncogenes and (iii) complex karyotypes (more than 3 chromosomal aberrations). Complex chromosomal aberrations (CCAs) are associated with the most unfavorable prognosis among subtypes of MDS and AML, and MDS cases with a complex karyotype have a high propensity to evolve to AML. Despite intensive treatment including allogeneic stem cell transplantation, long-term remissions are achieved in less than 10% of patients with CCAs. The frequency of CCAs is remarkably high: 20% of de novo AMLs, 30% of de novo MDSs, 24% of secondary AMLs and up to 50% of therapy related AML and MDS cases. The lower frequency of CCAs in de novo AML reflects the higher prevalence of classic translocation-generated oncogenes (e.g., AML1-ETO, PML-RARα and many others) in this disease compared with their paucity in MDS. Thus, myeloid leukemias and myelodysplastic syndromes with CCAs constitute important clinical entities in need of improved therapeutic strategies.

Cytogenetic studies in MDS have revealed primarily unbalanced chromosomal abnormalities and deletions, most commonly resulting in −5, 5q−, −7, 7q−, +8, 11q−, 13q− and 20q−, suggesting that genes within these regions have a role in MDS pathogenesis (Fig. 1). Finding the genes affected by such deletions poses a major investigative challenge, but will be necessary to accelerate progress in research and treatment of these myeloid diseases.

Clonal chromosomal abnormalities are observed in bone marrow cells from 30% to 50% of de novo MDS cases and 80% of secondary MDS patients. The predominant abnormalities discovered in MDS are nonrandom chromosomal deletions, suggesting a pathogenic mechanism based on loss of tumor suppressor genes or haploinsufficiency of genes necessary for normal myelopoiesis. Common cytogenetic abnormalities in MDS include loss of chromosome 7 or partial deletions of chromosome arms 5q, 20q, 11q, or 7q. In addition, juvenile chronic myelomonocytic leukemia often involves monosomy 7, together with mutations of the NF1 gene. In a study of 1663 cases of MDS, 1098 (66%) had a single chromosomal abnormality, 237 (22%) were monosomic for chromosome 7, and 431 (39%) had a partial deletion of chromosome 5. Other abnormalities included chromosomes 6, 9, 11, 12, 13, and 17. Importantly, cytogenetic abnormalities correlate with prognosis. After intensive chemotherapy, 60% of patients with an apparently normal karyotype entered complete remission (average duration, 16 months), while patients with chromosome 5 or 7 deletions or complex chromosomal abnormalities had a 20% remission rate (average duration, 4 to 5 months). Similarly, secondary MDS usually displays monosomies of chromosome 5 or 7 or partial deletions involving 5q or 7q, with chromosome 7 defects associated with decreased survival time.

Although the majority of putative tumor suppressors in MDS have not been cloned, many chromosomal translocation-mediated oncogenes (see Fig. 1, reviewed in Look3) and a few of the tumor suppressors have been identified. For example, genes inactivated in AML comprise a relatively small number of cases and include P53, RB, WT-1, NF1, AML1, C/EBPα, and nucleophosmin (NPM) (reviewed in Tenen6 and Side28), and besides p53, these genes are rarely mutated in MDS. In addition, the deleted region in the 5q− syndrome, which is commonly encountered in older individuals with MDS but uncommon in pediatric patients, contains the alpha-catenin gene. As we have recently shown, this gene is both deleted on one allele and epigenetically suppressed in a subset of cases with 5q−, most likely contributing to dysregulated and aberrantly increased stem cell self renewal within the clone.

MLL Amplification in MDS

In contrast to AMLs harboring oncogenic transcription factor fusions, hardly any oncogene activation has been assigned specifically to MDS and AML with CCAs. One exception is the 11q23 region and its resident MLL gene, which is amplified in a significant fraction of MDS and AML with karyotypic complexity and an adverse prognosis. MLL has long been recognized as an important component of translocation-generated fusion proteins. In contrast to other oncogenic fusion proteins, MLL participates in translocations with more than 40 different partner chromosomal loci. Homodimerization of the chimeric proteins appears to underlie the promiscuity of MLL in its ability to combine with many fusion partners, at least for a subset of its productive fusions.

The new studies mentioned above suggested that amplification of MLL represents a new mechanism of oncogenic activation of this gene. A recent study confirmed the importance of MLL within the 11q23 amplicon by analysis of MLL target genes.
like HOXA9 and MEIS1. It has long been recognized that HOXA9 is one of the important target genes of MLL and recent reports from several independent laboratories, including ours, provide a comprehensive view of other HOX genes that may be inappropriately activated in leukemias with MLL rearrangements. Very recently, Hoxa7 and Hoxa9 were shown to be essential for MLL-dependent leukemogenesis in vivo.

Thus, aberrant MLL activation in MDS/AML and consequent dysregulation of its downstream targets are of clinical importance. Indeed, the remarkable synergy between MLL gene amplification and loss of 5q in MDS and AML are reported to result in an extremely poor overall survival rate of 30 days. Moreover, these data provide an important clue regarding the mechanism of disease evolution. It should be stressed that the impact of dysregulated MLL and HOX gene activation is likely to extend beyond the subgroup of patients with CCA, as MLL and HOXA9 were also significantly upregulated in unselected MDS patient samples, including those with normal karyotypes.

**HOX Genes: Multiple Roles in Development, Hematopoiesis and Leukemogenesis**

Homeobox genes were first recognized through the analysis of homeotic mutations of Drosophila, which alter the identity of various body segments. Homologous genes have been found in virtually every species, from yeast to humans. Class I homeobox genes are designated as HOX genes in humans (mouse: hox genes). HOX genes control morphogenesis in early stages of embryonic development. The specific shape of discrete segments (pattern formation) is decisively regulated by these genes. Beside their role as differentiation factors in embryonic development, the control of hematopoiesis by HOX genes is well established. Since the perturbation of hematopoietic stem cell development is a hallmark of leukemia, it is not surprising that the aberrant expression of HOX genes contributes decisively to leukemia pathogenesis.

Early experimental evidence suggesting the oncogenic potential of the HOX gene family came from studies showing that the overexpression of Hoxb8 and IL-3 in murine bone marrow cells can induce aggressive, transplantable leukemia. A similar approach showed that Hoxa9 and Hoxa10 are able to induce AML in mice. Retroviral insertional mutagenesis has likewise implicated the Hox genes in leukemia induction. For example, Hoxa7 and Hoxa9 were activated in the context of retrovirally induced AML in BXH-2 mice. The importance of Hoxa7, Hoxa9 and the cofactor Meis1 in the BXH-2 mouse leukemia model was impressively underscored by a study based on large scale cloning of proviral integration sites.

Further compelling evidence of the oncogenic potential of HOX genes comes from their direct or indirect involvement in leukemia-associated translocations, such as the translocation t(7;11)(p15;p15), which generates the fusion protein NUP98-HOXA9 in AML patients. Additional translocations involving HOX loci have been identified over the past years. As pointed out above, MLL translocations (6%-7% of all acute leukemia) and most likely also MLL amplifications lead likewise to a dysregulation of HOX gene expression. Very recently CDX4 was shown to regulate expression of HOXA7 and HOXA9. Thus, this homeobox transcription factor and its relative CDX2 are attractive candidates for unidentified upstream regulators of HOX gene expression in leukemia. Taken together, published reports leave little doubt that specific HOX genes, particularly HOXA7, HOXA9 and HOXA10, are involved in the pathogenesis of AML and MDS, but virtually nothing is known about the downstream pathways through which these genes exert their oncogenic potential.

**References**


The Role of Allogeneic Stem Cell Transplantation in MDS-MPD

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Allogeneic SCT (alloSCT) is the treatment of choice for patients with MDS who have a histocompatible donor, but the question is up to which age? Patients with early stages of MDS may profit optimally from alloSCT, with long-term DFS in more than 50% of patients. Longer disease duration before transplantation is associated with an increased TRM after transplantation. A recent analysis confirmed that delayed transplantation may result in maximized overall survival for low risk MDS. They hypothesized that the optimal timing of transplantation for this cohort is at the time of development of a new cytogenetic abnormality, the appearance of a clinically important cytopenia or an increase in the percentage of marrow blasts. Whether patients with advanced stages of MDS should receive remission-induction chemotherapy prior to the transplant conditioning remains a point for discussion. Retrospective analyses showed conflicting data. Interpretation of the data is hampered by selection biases in the two treatment approaches. Therefore, the EBMT has launched a prospective study for this question (www.ebmt.org: Chronic leukemia; protocol: Allo-MDS 2x2). The results of alloSCT with unrelated donors have improved markedly in recent years.2 The American National Marrow Donor Program (NMDP) reported an improved DFS in more recently performed transplantations in a cohort of 510 patients with MDS transplanted with unrelated donors. The relative risk for DFS was 1.43 (95% confidence interval: 1.01–2.01) for transplantations performed between 1988 and 1993 versus more recent transplantations3.

The idea to reduce the intensity of the conditioning regimen has been developed in view of the high TRM of conventional marrow ablative conditioning regimens. The principle of RIC regimens is based on intensive immune suppression during conditioning and/or after stem cell infusion to facilitate donor engraftment and to establish complete donor chimerism. It is difficult to reconcile the contribution of RIC regimens to the improved outcome of alloSCT in view of the recently improved outcome of transplantation with marrow ablative regimens and the heterogeneity of the patient populations (age, co-morbidity, stage of disease).4 For this reason, the EBMT has launched a prospective randomized study comparing RIC regimens with standard conditioning regimens in patients with MDS older than 50 years.

The question remains whether autoSCT is a good alternative for alloSCT? In a recent study the 4-year DFS of the patients with a donor was 46%, higher than the DFS of 26% in patients without a donor. Subgroup analysis showed that the advantage of the presence of an HLA-identical sibling donor was only apparent in the patient group with intermediate and high risk cytogenetics.

General conclusion: allogeneic SCT may be considered the curative treatment option of choice. The upper age limit for alloSCT is mainly mandated by co-morbidity and general condition of the patient (frailty index) rather age of 60 or 70 years. For patients lacking an suitable donor, treatment with autoSCT or chemotherapy may be a good alternative for MDS patients with good-risk cytogenetic characteristics.
References


Multiple Myeloma (MM) is a chemoresistant malignancy and for decades, the only active drugs were alkylating agents and high-dose corticosteroids. The introduction of novel agents (Thalidomide, Bortezomib and Lenalidomide) is changing the management of MM patients both for frontline therapy and at relapse. These agents have a different mode of action and can act not only on the myeloma cells but also on the microenvironment which is necessary for tumor cell survival and proliferation. In vitro experiments have shown possible synergy with dexamethasone or chemotherapeutic agents. They are active even in heavily pretreated patients and offer new possibilities for relapsed/refractory MM.

**Thalidomide**

*Thalidomide as a single agent*

Thalidomide is administered orally. Due to its teratogenicity, it is prescribed according to risk-management programs to avoid exposure of women with child-bearing potential.

Thalidomide was introduced in the treatment of relapsed MM by the Arkansas group (1). Following the results of this pioneer work, a large number of Phase II studies were reported and were summarized in a comprehensive review (42 studies, 1674 patients) (2). From this experience, the following conclusions are possible (2-5).

- The response rate is approximately 45% including 30% partial remissions (PR)
- Complete remission (CR) are rare in heavily pretreated patients but possible
- The onset of response is rapid (less than 2 months) and maximal response is achieved within 6 months
- One-year event-free survival (EFS) is about 35% and median overall survival (OS) is 14 months
- Prognostic factors with thalidomide treatment are the same as with conventional chemotherapy and include tumor burden markers and cytogenetic abnormalities
- Thalidomide can be prescribed in patients with renal dysfunction
- Myelosuppression is very rare
- The incidence and severity of side effects is related to the daily dosage. With the doses initially used (400 mg/day) the most frequent toxicities were constipation, somnolence, fatigue and peripheral neuropathy.

Peripheral neuropathy causes numbness, paresthesia and even pain in legs and arms. The overall incidence of peripheral neuropathy is 30% with 10% grade ≥ 3, but is related to cumulated dose and is up to 75% in patients treated more than one year (6). Since there is no effective prophylaxis and treatment, the drug should be discontinued in case of peripheral neuropathy signs or symptoms (3-5).

Currently the daily doses of Thalidomide have been reduced to 100-200 mg/day.

**Thalidomide in combination**

The combination of Thalidomide plus Dexamethasone has been developed with the objectives of increasing the efficacy and of reducing the daily dose and the toxicity of Thalidomide (6-8). Although...
no randomized study comparing Thalidomide and Thalidomide/Dexamethasone (TD) has been performed, TD is considered more effective than Thalidomide alone (approximatively 45% response rate) and superior to conventional chemotherapy as treatment of first relapse (9). Since Thalidomide is not myelotoxic, combinations with chemotherapy have also been evaluated in Phase II trials with response rates ranging from 36% to 73%.

While combinations of Thalidomide with Dexamethasone or chemotherapy alone appear to be more active, they are more toxic with a higher incidence of infectious complications. Most importantly they induce an unexpected complication, deep vein thrombosis. While with Thalidomide alone the incidence of deep vein thrombosis is < 5% as with any treatment of MM, with TD the incidence is 10-15% and with chemotherapy (specially anthracyclines) it increases up to 30% (10, 11). This complication usually occurs during the first 3 months of treatment and is more frequent in newly diagnosed patients and in patients with a high tumor burden. The optimal prophylaxis is not yet known and oral anticoagulant, low molecular weigh heparin and low-dose aspirin are currently evaluated (3).

**Bortezomib**

Bortezomib is the first in class proteasome inhibitor. In MM it is active not only on the myeloma cell but also on the microenvironment which is necessary for tumor cell proliferation and survival. It is administered IV at a dose of 1.3 mg/m² on days 1, 4, 8, 11 in 21 days cycles. Two Phase II studies have shown that in heavily pretreated patients, response rate with Bortezomib as a single agent is 25-30% and can be increased by the addition of Dexamethasone (12, 13). Following these studies, the drug has been approved in the US and Europe. The large randomised Phase III trial APEX has demonstrated that Bortezomib is superior to Dexamethasone in relapsed MM, in terms of response rate (including CR), time to progression and OS (14). The onset of response is rapid, usually within 2 cycles but the maximal response can be achieved after up to 8 cycles. A subgroup analysis on patients having received one line of treatment confirmed the superiority of Bortezomib compared to Dexamethasone as treatment of first relapse (15). Based on these Phase II-III studies the toxicity profile is well defined. The most frequent side effects are gastrointestinal symptoms (diarhea or constipation) and fatigue but they are usually mild. The drug is not myelotoxic but can induce up to 60% decrease of the platelet count (16). This thrombocytopenia is rapidly reversible usually before the following cycle. Peripheral neuropathy is observed in 30-40% of cases (grade ≥ 3 in 10-15%). Signs and symptoms are reversible in 2/3 of cases after dose reduction or drug discontinuation (17). Bortezomib appears to be as effective in older patients and in patients with poor risk disease (18, 19). It can be prescribed safely in patients with renal failure, even in patients on dialysis (20, 21). Based on preclinical studies and on the drug toxicity profile, a number of Phase II studies have evaluated Bortezomib in combination with either Dexamethasone or with chemotherapy.

**Lenalidomide**

Lenalidomide is a Thalidomide analog which appears to be more potent in vitro. It is not teratogenic in animal models but most importantly, after the Phase I study it became apparent that the toxicity profile of Lenalidomide was completely different (22). Constipation, somnolence, fatigue and peripheral neuropathy that are frequent adverse events with Thalidomide, were not observed. The most frequent side effect was myelosuppression mostly after 28 days of treatment. In Phase II trials the drug was given orally for 21 consecutive days. The response rate in heavily pretreated patients was 25% and was increased by the addition of Dexamethasone (23). Two large randomized trials were then conducted in the US and in Europe (24, 25). They both compared Lenalidomide (25 mg/day on 21 consecutive days) plus high-dose Dexamethasone versus Dexamethasone. They both showed the superiority of the combination in terms of response rate (including CR), time to progression and OS. Based on these studies, the drug was approved by FDA in 2006 and the European approval is pending. However, like with Thalidomide, the combination of Lenalidomide with Dexamethasone induces deep vein thrombosis and justify prophylactic treatment at least with low-dose aspirin. The treatment should be used with caution in patients with renal failure due to a higher myelotoxicity (specially thrombocytopenia) in patients with less than 50 ml/min creatinine clearance. Lenalidomide is currently tested upfront in combination with dexamethasone or with chemotherapy.

**Combination of novel agents**

Since the toxicity profiled of these three agents is different, it appeared logical to combine them
with the objective of increasing efficacy. The combination of Thalidomide or Lenalidomide with Bortezomib was the most attractive, due to a possible synergy of agents having different mode of actions. Thalidomide and Bortezomib have been combined with either Dexamethasone (VTD) or with Melphalan Prednisone (VMPT) with very encouraging results and an acceptable toxicity (26, 27). The combination Bortezomib-Lenalidomide is currently tested (28).

**Role of Stem Cell Transplantation at first relapse**

Autologous Stem Cell Transplantation (ASCT) is currently considered the standard of care for frontline therapy in patients up to 65 years of age. However ASCT is also a useful salvage treatment in chemosensitive or untreated relapses (29). When comparing early ASCT, and ASCT when conventional chemotherapy fails (late ASCT) there is no difference in OS although time to progression is longer with early ASCT (30). If late ASCT is considered it should be useful to collect stem cells early (31) since the hematopoietic quality of grafts is often decreased by previous chemotherapy specially alkylating agents (32). When ASCT has been performed upfront, a second ASCT is justified if the duration of first remission has been > 2 years.

Allogeneic SCT is probably the only treatment that can induce long term clinical and molecular remissions. However standard myeloablative regimens prior to allogeneic SCT have been almost abandoned specially for relapsed MM, due to a high transplant-related mortality (up to 50%) (33).

With reduced-intensity conditioning allogeneic SCT, transplant-related mortality is reduced, even in relapsed MM (20-25% at 1 year) (34). However the risk of relapse is higher than with standard allogeneic SCT, specially in chemoresistant disease or when the tumor burden is high (35). Current strategy aims at reducing the tumor burden first, for instance with high-dose Melphalan followed by ASCT, and then to exploit the graft-versus-myeloma effect of donor lymphoid cells. This strategy is mostly proposed for first line treatment in patients with an HLA-identical donor (36) but can also be offered at first relapse if CR or VGPR has been achieved with salvage therapy.

**Therapeutic strategy**

With the introduction of novel agents, CR achievement is possible in relapsed MM. In the large randomized studies testing Bortezomib and Lenalidomide, the progression-free survival (PFS) and OS increase was associated to a CR rate increase, as compared to the control arms (12, 24, 25). In the APEX trial, PFS was 12 months for patients in CR versus 8 months for patients with only PR (37). Therefore, as for frontline therapy, the objective of first relapse treatment should be to achieve the best possible response.

With this objective, it is logical to use novel agents for first relapse treatment since they are effective even in heavily pretreated patients. Although no randomized study has compared Thalidomide, Bortezomib or Lenalidomide given as single agents to combinations, they are usually administered in combination at least with dexamethasone (TD, VD or RD). Out of a clinical trial, the choice depends on a variety of parameters including age, previous treatment, prior toxicities, availability of novel agents (which is not the same all over the world). There are special situations in which the choice is easier.

For instance if Thalidomide has been used for frontline therapy it is better to use Bortezomib or Lenalidomide to avoid the risk of peripheral neuropathy related to cumulative doses of Thalidomide. In patients with renal failure, Bortezomib is the treatment of choice while in patients with peripheral neuropathy, Lenalidomide should be preferred. Fulminant relapses or poor risk cytogentics should be treted with combinations including Bortezomib or Lenalidomide.

The optimal duration of salvage treatment is unknown. In order to reduce the risk of toxicity and of resistant clones selection, treatment could be stopped 3 months after best result is achieved. In younger patients (< 65 years) ACHS can be considered in sensitive relapses if stem cells are available. Reduced intensity conditioning allogeneic SCT is possible in patients who achieve CR with salvage treatment.

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Reduced Intensity Conditioning (RIC)
Allogeneic Stem Cell Transplantation for LLM: Hype, Reality or Time for a Rethink

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Abstract
Reduced-intensity conditioning has been increasingly used before allogeneic SCT in a growing number of indications. It is well established that the first goal, allowing SCT for elderly and medically infirm patients has been achieved. RIC regimens result in consistent engraftment of allografts from related and unrelated donors. TRM rates have been markedly reduced, thus SCT can be administered relatively safely with no upper age limit, after prior autologous SCT, and in certain malignancies, such as lymphomas and myelomas, in which TRM rates were exceedingly high. However, toxicity may still be substantial with some regimens and in patients with a high comorbidity score. GVHD continues to be a major cause of morbidity and mortality after RIC and its incidence may not be lower than after myeloablative SCT. Invasive fungal infections are a second common cause of TRM, which are closely associated with GVHD, did not reduce in incidence. Novel approaches to further, reduce these two complications are required to further improve outcome. In-vivo T-cell depletion reduces initial rates of GVHD, but because DLI is required more often for increased risk of disease persistence and MC, the ultimate rate of GVHD remains unchanged. With gained experience, DLI is used more carefully after RIC, limiting is use to patients with persistent MRD after SCT or imminent graft rejection and delaying administration in other patients, may reduce the risk for GVHD. Methods to deliver cellular immune therapy without GVHD would be a major step forward. The development of tumor or minor histocompatibility antigen-restricted DLI and the combination with targeted therapy and tumor vaccines are promising. Despite initial data, there is no firm evidence for advantage of any of the regimens over the other regimens. Although TRM may reduce, with RIC, relapse rates may increase compared with ablative SCT, thus the net effects on disease-free survival are yet to be determined. Prospective comparative studies to determine the best NST regimen and randomized studies comparing RIC and ablative SCT are urgently required before RIC can be accepted as standard therapy and to better define its role.

Background
Allogeneic hematopoietic stem cell transplantation (SCT) is an effective potentially curative treatment of advanced or high-risk hematologic malignancies. High-dose chemotherapeutic regimens is a major step toward achieving this goal. Much experience has been gained with this approach over the past decade.

Rationale for RIC Stem Cell Transplantation
Stem cell transplantation was initially developed as a means to deliver high-dose chemotherapy and radiation for elimination of the underlying disorder. Escalation of treatment doses results in better tumor kill, but it leads to irreversible myelo-suppression. SCT was viewed as a supportive-care modality to restore hematopoiesis after treatment. However, it has subsequently become apparent that high-dose chemoradiotherapy does not eradicate the disease in many patients and that much of the therapeutic benefit of SCT relates to an associated, immune-mediated, graft-versus-leukemia (GVL), or graft-versus-malignancy (GVM) effect, which led to a novel therapeutic approach. Low-dose relatively nontoxic and tolerable conditioning regimens have been designed, not to eradicate...
the malignancy, but rather to provide sufficient immunosuppression to achieve donor cell engraftment and to allow induction of GVL as the primary treatment.

RIC stem cell transplantation does not eliminate all host hematopoiesis and commonly leads to a state of mixed chimerism (MC). MC describes persistence of donor cells with normal host hematopoietic cells and cells of the underlying malignancy. Stable long lived MC has been reported in animal models and in patients having RIC SCT for nonmalignant disorders. However, in patients with malignancies, MC is most often transient and conversion to complete chimerism (CC), autologous recoRIC SCT, or relapse occurs spontaneously or after immune manipulations within the first few months after RIC SCT. The initial nonmyeloablative treatment is expected to produce only transient suppression of the underlying malignancy, but it allows time for the immune GVM effect to develop. Patients with MC or with detected or minimal residual disease (MRD) after RIC SCT may require additional immune-therapeutic approaches. Immunosuppressive therapy administered after SCT for the prevention of GVHD can also suppress GVL. Early withdrawal of immunosuppressive therapy allows the occurrence of potent GVL effect that can potentially eliminate residual disease and host hematopoiesis producing complete remission and CC, respectively. If this does not occur, donor lymphocyte infusions (DLI) may harness this effect and switch the balance toward CC/complete remission. The GVL and graft-versus-hematopoietic tissue effects are highly associated with GVHD, although they may also occur in its absence. Therefore, the initial RIC SCT serves as a platform for additional allogeneic cellular therapy.

RIC Stem Cell Transplantation Regimens

Nonmyeloablative stem cell transplantation regimens comprise of a spectrum of regimens with different immunosuppressive and myelosuppressive properties. The kinetics of engraftment, chimerism, and eradication of residual disease differ accordingly. Conditioning regimens have been referred to as nonmyeloablative if they do not completely eradicate host hematopoiesis and immunity. A few of these regimens have been administered as chemotherapeutic regimens with no stem cell support and allow relatively prompt hematologic recovery. Autologous recoRIC SCT itution of hematopoiesis is expected if the allograft is rejected. These regimens are only mildly myelosuppressive and commonly result in induction of MC. CC and GVL may develop slowly, spontaneously, or after immune interventions. The Seattle regimen consisting of low-dose total body irradiation (TBI: 200 cGy) with (or initially without) fludarabine and intensive pre and post transplant immunosuppression is the prototype of these regimens.

More intensive regimens have also been developed. These regimens have been referred to as reduced intensity conditioning regimens. They have not been administered without stem cell support and autologous recovery may be slow, if at all. These regimens often combine immunosuppressive agents, such as fludarabine with or without sero-therapy (antithymocyte globulin [ATG] or alemtuzumab), and agents with moderate myelosuppressive effects, such as busulfan or melphalan. Although these regimens are more intensive than the nonmyeloablative regimens, dose intensity is still reduced compared to ablative regimens, allowing reduction of toxicity. Reduced intensity regimens, in similarity to myeloablative regimens, rapidly induce CC and antitumor responses, but they are more toxic and are associated with a higher risk for GVHD. A third approach is using a double-step strategy. High-dose chemotherapy supported by autologous SCT is used for cytoreduction and also as an immunosuppressive platform for the second stage of allogeneic SCT with nonmyeloablative or reduced-intensity conditioning administered 2 to 3 months later. The separation of high-dose chemotherapy and allogeneic effects results in reduced toxicity and better tolerability compared to when allogeneic SCT immediately follows high-dose chemotherapy. A novel approach is to combine RIC SCT with targeted therapy. Imatinib mesylate has been explored as adjuvant to RIC SCT before SCT, allowing reduction of conditioning intensity, and after SCT to eliminate MRD. Rituximab has been used in conjunction with RIC SCT in lymphoid malignancies and by us after SCT to target MRD. Radio-labeled immune conjugates, such as radio-labeled anti-CD20 monoclonal antibodies, may be used with SCT to target lymphoma cells, allowing the use of less intensive conditioning.

RIC Stem Cell Transplantation and Regimen-related Complications

Nonmyeloablative stem cell transplantation regimens were originally designed to enable treatment of older and medically infirm patients not eligible
for ablative conditioning. This goal has largely been achieved. Standard ablative regimens are often limited to patients up to age 55 years. Age was not found to be an adverse factor for prediction of outcome after related and unrelated donor RIC SCT and is no longer a contraindication for SCT. Standard SCT in certain high-risk settings, such as in patients failing a prior autologous SCT, and in patients with certain diagnoses, such as multiple myeloma, Hodgkin’s, and non-Hodgkin’s lymphoma, was associated with unacceptably high treatment-related mortality (TRM) rates as high as 50%. TRM in the range of 10% to 20% can be observed in these settings using RIC SCT regimens. In particular, RIC SCT was able to reduce TRM after unrelated donor SCT.

Reduction of TRM is largely attributed to reduction in organ toxicity. The Seattle group has shown marked reduction in cardiovascular, gastrointestinal, hepatic, infectious, metabolic, neurologic, and pulmonary toxicity when comparing their low-dose TBI-based nonmyeloablative regimen to ablative regimens. Nonrelapse mortality within the first 100 days was 9% and 21%, respectively. The major therapy-related organ dysfunction syndromes are reduced in incidence. In particular, idiopathic pneumonia syndrome is less frequent after RIC SCT (2.2% vs 8.4% in one study), despite treatment of older patients. Hepatic toxicity may still be substantial, especially after some reduced intensity regimens. However, not all syndromes are reduced. We have shown that thrombotic microangiopathy is a frequent devastating complication after RIC SCT, more common in second SCTs and in association with acute GVHD. Diffuse alveolar hemorrhage is also relatively common in this setting. We have hypothesized based on experimental data that fludarabine-related endothelial and pulmonary epithelial toxicity may be associated with this unexpected observation. Other hematologic complications associated with donor-recipient ABO incompatibility may be more common after RIC SCT. Although direct toxicities of high-dose chemotherapy are reduced with RIC SCT, toxicities involving immune mechanisms may not. Organ toxicities are largely associated with patient comorbidity score before SCT. Further research is required to define the relative organ toxicities in different regimens.

RIC stem cell transplantation is less myelosuppressive compared to ablative conditioning: which results in a shorter duration of neutropenia and less transfusion requirements. Some of the nonmyeloablative regimens result in minimal myelosuppression and can be safely administered in the outpatient setting. Reduced-intensity regimens often result in more profound cytopenias more similar to ablative conditioning. The reduced duration of neutropenia and the limitation of mucosal injury result in reduced risk for severe infections in the immediate post-SCT period. However, the risk for invasive fungal infections is not reduced. These infections are often associated with GVHD and corticosteroid therapy and represent one of the major causes of TRM after NSI. In the Seattle study, invasive fungal infections occurred in 19% of RIC SCT recipients, relatively late in the course, and were the primary cause for 39% of nonrelapse-associated deaths.

**RIC Stem Cell Transplantation and Graft-versus-host Disease**

Graft-versus-host disease is one of the major causes of post-SCT morbidity and mortality. When RIC SCT was introduced, it was hoped that GVHD incidence would reduce. Acute GVHD results at least partially from tissue injury and cytokine release secondary to the toxicity of the preparative regimen, amplified by donor immune cells. The use of RIC SCT should theoretically limit tissue injury and cytokine release and reduce the incidence and severity of GVHD. MC that is more common after RIC SCT allows bilateral transplantation tolerance with some protection from GVHD. However, host-antigen presenting cells that have a major role in initiation of GVHD may persist after RIC SCT and contribute to GVHD. The duration of immunosuppressive therapy is usually shorter after RIC SCT and immune manipulations are often incorporated into RIC SCT programs, increasing the likelihood of GVHD, although delayed immune manipulations, once the toxicity of conditioning and cytokine release are already resolved, are less likely to produce severe GVHD. The net effect of these differences between RIC and ablative SCT on GVHD is still not well established. The Seattle group reported that the incidence of grade II/IV acute GVHD after RIC SCT was significantly lower than after ablative therapy, reaching 64% and 85%, respectively. However, the incidence of chronic GVHD was approximately 70% in both cohorts. Initiation of steroid therapy was delayed from an average of 1 to 3 months after SCT, corresponding to a new syndrome described as late-onset acute GVHD. This study suggests that GVHD is not reduced in incidence with RIC SCT, but it is only delayed. In
another study, The incidence of grade II/IV acute GVHD of 36% after myeloablative regimens, but only 12% after truly nonmyeloablative regimens. Chronic GVHD was also reduced. Further prospective studies are needed to determine the relative incidence of GVHD after RIC SCT. However, because it is still a major cause of morbidity and mortality, several approaches have been explored to decrease the risk.

Initially, RIC SCT regimens called for only a short course of immune suppression and early administration of DLI for disease eradication and conversion to CC. These interventions markedly increase the risk of GVHD. More recently, more careful approaches were introduced including the extension of the duration of immune suppression, especially after unrelated donor transplantation, up to 6 months. With better understanding of chimerism and MRD kinetics, the indications for DLI have been restricted, trying to reserve it only for patients destined to relapse or reject their graft, thus reducing the risk of GVHD in all other patients.

Another approach is the use of in-vivo T-cell depletion. Alemtuzumab is an effective agent in the prevention of GVHD. Alemtuzumab administered during conditioning depletes host T cells, thus reducing the risk of graft rejection reported with in-vitro T-cell depletion techniques. Alemtuzumab also depletes host-antigen presenting cells involved in GVHD. It persists after SCT and partially depletes donor T cells, thus alemtuzumab is very effective in the prevention of GVHD. However, patients administered alemtuzumab have a higher risk of opportunistic infections, in particular cytomegalovirus. Furthermore, alemtuzumab recipients have a higher risk of MC and residual disease and require more DLI, thus after DLI, the ultimate net risk of GVHD is not reduced and there is no improvement in survival. ATG administered before SCT has similar effects, although it may be less effective in the prevention of GVHD. Studies are being conducted to determine the dose of alemtuzumab or ATG that may result in net effects that would improve survival.

**Immune-therapeutic Intervention After RIC Stem Cell Transplantation**

In indolent malignancies, such as chronic myeloid leukemia (CML), chronic lymphocytic leukemia (CLL), follicular lymphoma, and multiple myeloma, MRD can be followed and no intervention is indicated unless progression or plateau in response is observed or quantitative MRD is rising. In aggressive malignancies, such as acute leukemia, timing is more crucial. There may not be sufficient time to follow MRD because relapse may occur within weeks, whereas effective DLI response may take 2 to 3 months. The sensitivity of the test is important. DLI may be administered early or when using very sensitive tests, such as quantitative polymerase chain reaction, when applicable. MRD can be followed very closely (eg, every 1-2 weeks). If MRD is declining, no intervention is needed.

The kinetics of MRD after RIC SCT is not well established as after ablative conditioning. The same level of MRD, may not necessarily have the same significance. MRD surviving high-dose chemotherapy and, to a lesser extent, reduced-intensity conditioning represents highly resistant malignancy, whereas MRD is expected after RIC SCT. MRD remaining after T-cell depletion or the use of alemtuzumab is also highly predictive of relapse.

In the future, tumor-specific lymphocytes or DLI generated against hematopoietic-specific minor histocompatibility antigens, such as HA-1 and HA-2, may be used to harness anti-tumor responses without the risk of GVHD and follow SCT with T-cell depleted grafts.

Targeted therapy is another option for control of MRD. Imatinib mesylate may be effective in salvaging CML patients with relapse or persistent disease after SCT, frontline or after failure of DLI. Imatinib mesylate may be synergistic with DLI. We have shown that rituximab administered after SCT for aggressive lymphoma reduced relapse risk in very high-risk patients. Rituximab may have eliminated MRD and may have synergized with the donor immune system, providing effectors for antibody-dependent cytotoxicity. Similar effects of rituximab administered for residual CLL after RIC SCT. Future studies may identify other methods to target MRD, trying to reduce relapse risk after SCT.

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Hairy cell leukemia (HCL) is a B-cell chronic lymphoproliferative disorder characterized by splenomegaly, pancytopenia, circulating lymphocytes displaying prominent cytoplasmic projections and reactive marrow fibrosis [1, 2]. HCL usually has an indolent disease course, and similarly to chronic lymphocytic leukemia, patients who are asymptomatic do not require therapy. However, treatment should be considered for symptomatic or cytopenic patients. Treatment is indicated for patients with significant neutropenia, thrombocytopenia, symptomatic splenomegaly, constitutional symptoms due to HCL or recurrent serious infections [3].

The first active agent for this disease was interferon-α (IFN-α) which can produce a complete response (CR) in approximately 10% of HCL patients and a partial response (PR) in the majority of remaining patients [4-7]. More recently, new purine nucleoside analogs (PNA), cladribine (2-chlorodeoxyadenosine, 2-CdA) and pentostatin (deoxycoformycin, DCF) have become a gold standard in the treatment of this disease [8-11]. 2-CdA and DCF are the drugs of choice in the treatment of HCL. Both agents are equally active, and have impressive long-term effectiveness [5]. However, monoclonal antibody (MoAb) directed against CD209 and immunotoxins directed against CD22 or CD25 have been introduced and are currently a promising novel approach to the treatment of resistant HCL [12-14].

Cladribine

Multiple studies demonstrated that 2-CdA induces durable and unmaintained CR in about 80% of patients after a single course of therapy [15-23]. 2-CdA is administered either as a continuous i.v. infusion at a dose of 0.09 mg/kg over a 5-7 days period or as 2-hour i.v. infusion at a dose of 0.12 mg/kg for 5-7 days. However, similar results were achieved when the drug was given in subcutaneous injection [19,24]. Patients in apparent clinical and hematological remission following a single course of 2-CdA administration may have residual disease detected with the use of flow cytometry or molecular assay. 2-CdA is also highly effective in relapsed disease. Goodman et al. [20] has reported the results of re-treatment with this agent. The overall response rate of relapsed patients who were retreated with 2-CdA was 90% with 75% achieving a CR and 10 (17%) a PR. The median second response duration for all responders was 35 months while the median time to first relapse was 42 months. The second relapse was observed in 20 patients and 10 of them received second re-treatment with 2-CdA. Overall response rate was 80% including 60% CR. The median third response duration was 20 months.

2-CdA is also an effective drug when administered at a dose of 0.15 mg/kg in 2-hour infusion once a week over 6 course [22]. In our randomized study we compared weekly administration of 2-CdA (0.12 mg/kg in 2-h i.v. infusion once a week for 6 weeks) with daily administration (0.12 mg/kg in 2-h i.v. infusion for 5 consecutive days) [23]. The final results of this study indicate that both CR and overall response (OR) rates were similar in compared groups. There was no statistically significant difference in toxicity between groups. Moreover, HCL treatment with weekly 2-CdA infusion is not safer than standard 5-day 2-CdA.

Pentostatin

DCF is also a highly active agent in HCL [25-27]. The drug was usually used at a dose of 2-4
mg/m² i.v. every second week although other schemes were also applied. Ranges of CR varied from 40 to 90% regardless of prior treatment with IFN-α or splenectomy. The drug is well tolerated in HCL, especially when neutropenia is not severe or there is no history of life-threatening infections. Myelosuppression is the main toxicity and may require delays in planned chemotherapy schedule.

A randomized study comparing DCF with IFN-α has demonstrated that DCF produced a higher CR and PR rate with more durable responses in HCL [7]. In that study patients were randomized to receive either IFN-α (3x10⁶/l subcutaneously 3 times per week) or DCF (4 mg/m² i.v. every 2 weeks). Patients who did not respond to initial treatment were crossed over. Among IFN-α patients, 17 of 159 (11%) achieved a confirmed CR or PR. Among DCF patients, 117 of 154 (76%) achieved a confirmed CR and 121 of 154 (78%) had a confirmed CR or PR. Response rates were significantly higher (p < 0.0001) and relapse free survival was significantly longer with DCF than IFN-α (p < 0.0001).

Subsequently, long-term data on duration of overall survival and relapse-free survival data from this study were reported [27]. Estimated 5- and 10-year survival rates for all patients were 90% and 81%, respectively. Moreover, only 2 of 40 death were attributed to HCL. Other long-term observations also confirmed that DCF induces a high rate of long-lasting CR both in patients previously untreated and in patients pretreated with IFN-α or splenectomy [26,29].

2-CdA and DCF seem to induce similar high response rates but only a large randomized trial with the two agents will be able to evaluate the CR rates, duration of response, recurrence rates and adverse events, that have appeared to be comparable so far.

Recently, Else et al. [29] have reviewed a series of 219 patients with HCL, with median follow-up from the diagnosis of 12.5 years (range 1.0-34.6 yrs) treated either with DCF (n=185) or 2-CdA (n=34), to compare the effectiveness of these agents. Overall response to DCF was 96% with a CR in 81% and median disease free survival (DFS) of 15 years. Response to first line 2-CdA was 100% with a CR in 82% and DFS of 11+ years. DFS showed no plateau in both groups. The relapse rates at 5 years and 10 years were 24% and 42%, respectively, with DCF and 33% and 48% with 2-CdA. Survival at 10 years was 96% in DCF group and 100% in 2-CdA group. This study has confirmed previous observations that DCF and 2-CdA are equivalent in the treatment of HCL. However, 2-CdA affords the convenience of a single course of administration. Patients treated with PNA do relapse and the overall survival curves have not reached a plateau, which indicates that cure has not been secured.

**Rituximab and immunotoxins**

HCL presents the highest percentage of CD20 expression of lymphoproliferative disorders. Recently, rituximab as well as anti-CD22 and anti-CD25 immunotoxins have been investigated in refractory and/or relapsed HCL [5,30]. The use of MoAb therapy for the treatment of HCL offers also great promise and potential for improving progression free survival [42-57].

Lauria et al. [12] treated patients with HCL, who had progressed after prior therapy with 2-CdA and/or DCF, with rituximab. All patients received 375 mg/m² i.v. of rituximab once a week for 4 dose. Overall, out of 10 treated patients one achieved a CR, 4 - a PR and 3 - a minor response..

Twenty-four HCL patients reported recently by Nieva et al. [31] were treated with standard doses of rituximab. All of them relapsed after prior treatment with 2-CdA. Six patients achieved a response following rituximab including 3 (13%) with a CR and 3 (13%) with a PR. At a median follow-up of

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Fig 1. Proposed treatment algorithm for hairy cell leukemia (modified from [8]).
14.6 months, 2 responders relapsed. Even better result were presented by Thomas et al. [32]. Fifteen patients with relapsed or primary refractory HCL after 2-CdA or DCF received rituximab 375 mg/m² weekly at a total of 8 planned doses. The overall response rate was 80% including 53% CR.

Rituximab seems to be an ideal drug for elimination of MRD in HCL patients after treatment with PNA. Cervetti et al. [33] evaluated a role of this antibody in 10 patients previously treated with 2-CdA. Before starting rituximab, two patients were in CR, six in PR and two showed no significant response to 2-CdA. Rituximab was infused once a week at a dose of 375 mg/m² for four doses. Two months after the end of rituximab therapy, all evaluated patients were in hematological CR. Moreover rituximab increased percentage of molecular remission up to 100% one year after the end of treatment. These results have been recently confirmed by Ravandi et al. [14]. They have treated 13 patients, 11 with newly diagnosed disease and two after failure of one prior therapy with 2-CdA at a dose of 5.6 mg/m² daily for 5 days followed by rituximab at a dose of 375 mg/m² weekly for 8 weeks. All patients achieved a CR, and MRD was eradicated in 12 (92%) of the patients.

Promising results in patients with resistance to purine analogues have been obtained with two immunotoxins BL22 and LMB-2 targeting CD22 and CD25, respectively [13,34,35]. Recently, Kreitman et al. [13] have reported the results of their study testing the safety and efficacy of recombinant immunotoxin containing anti-CD22 variable domain (Fv) fused to truncated Pseudomonas exotoxin (RF B4 (dsFv)-PE-38) named BL22. Sixteen HCL patients who were resistant to 2-CdA were included into the study. All patients had circulating hairy cells that expressed CD22. BL22 at doses between 0.2-4.0 mg was administered as a 30 min intravenous infusion every other day to a total of 3 doses. Of 16 patients treated with BL22, 11 (69%) had a CR and 2 had a PR. Common toxic effect was transient hypoalbumina and the elevated aminotransferase level. Two patients developed reversible hemolytic uremic syndrome.

**Current indications for splenectomy and interferon-α**

With the introduction of PNA, the indication for splenectomy is extremely limited. The removal of the spleen may be appropriate in patients with resistant massive symptomatic splenomegaly or results in severe cytopenia following PNA therapy [60], and when progressive HCL developed during pregnancy [36,37]. Splenectomy may be also required for diagnostic reasons in patients with primary splenic HCL and in rare patients refractory to PNA, IFN-α and MoAb. Splenic rupture is also rare indication for splenectomy [58].

IFN-α may still have a place in HCL in pregnancy [64] and in the patients presenting with very severe neutropenia to increase neutrophil count prior to PNA therapy [38,39]. There is no significant role for IFN-α in improving the proportion and the duration of CR in HCL patients previously treated with DCF. In the study performed by Marotta et al [40] 135 patients who obtained CR or PR after treatment with DCF were randomized to receive IFN-α or not. Progression of disease was observed in 8 and 12 patients, with a median time of 27.8 and 26.9 months, respectively.

**Conclusions**

Treatment of progressive symptomatic HCL includes a variety of pharmacological approaches such as INF-α, DCF and 2-CdA, which have significantly improved the disease prognosis. 2-CdA and DCF seem to induce a similar high response rate and a long overall survival. They are also active in relapsed patients. About 80% of patients treated with 2-CdA or DCF survive for at least 10 years. In patients, who failed PNA therapy, anti-CD20 MoAb rituximab and anti-CD22 (BL22) immunotoxin appear extremely effective. Future studies should be directed to optimizing the therapy for minimal residual disease as well as whether the introduction of new effective agents will translate into cure. A proposed flow-chart for therapeutic decision-making in the treatment of HCL is outlined in Fig 1.

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**References**


First Line Treatment of Multiple Myeloma

Guido Tricot

University of Arkansas for Medical Sciences, USA

Tenets

- Tumor Stem Cell - Elusive
  - MM germinal center-derived post-switch stage
- Homing and Expansion in Bone Marrow (MM Strong Cross-Talk)
  - Exploiting the microenvironment including OB/OC for survival: new target for therapy

Multi-Step Progression of Multiple Myeloma

Cancer Stem Cells

- Non-cycling
- Excellent detoxification

Growth and Survival Control of Myeloma

- Non-cycling
- Excellent detoxification
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1Masood Bazrgar, 2Mehran Karimi, 2Masoumeh Talebi, 2Mahin Farahmand Beigi
1Human Genetic Research Group, Iranian Academic Center For Education, Culture & Research (acecr), Fars Province Branch, Shiraz, Iran, 2Hematology Research Center, Shiraz University of Medical Sciences, Shiraz, Iran.

LATE EXTRAMEDULLARY RELAPSES OF ACUTE LEUKEMIA
1Nesrin Karabul, 1Salmai Turial, 1Peter Gutjahr
1Childrens Hospital of University Mainz, Department of Pediatric Oncology, Germany 2Childrens Hospital of University Mainz, Department of Pediatric Surgery, Germany

RESULTS OF HYPERFRACTIONATED CYCLOPHOSPHAMIDE, VINCRISTINE, DOXORUBICIN, AND DEXAMETHASONE (HYPER-CVAD) IN ADULT ACUTE LYMPHOBLASTIC LEUKEMIA: SINGLE CENTER EXPERIENCE
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METHOTREXATE INDUCED NEUROTOXICITY DURING HYPER CVAD TREATMENT IN A PATIENT WITH ALL
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METHOTREXATE INDUCED NEUROTOXICITY DURING HYPER CVAD TREATMENT IN A PATIENT WITH ALL
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Ondokuz Mayis University, Samsun, Turkey
ABBERANT ANTIGEN EXPRESSION IN 236 PATIENTS WITH ACUTE MYELOID LEUKEMIA BY MULTI-COLOR FLOW CYTOMETRY

Mesude Yilmaz, Gulsum Ozet, Simten Dagdas, Funda Ceren, Meltem Ayli, Osman Yokus, Ozlem Balcilik, Murat Albayrak, Ayla Gokmen Aköz, Zeynep Aki, Ankara Numune Education And Research Hospital, Ankara, Turkey

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CD33–DIRECTED THERAPY WITH GEMTUCUMAB OZOGAMICIN IN ACUTE MYELOID LEUKEMIA: REPORT OF TWO CASES

Ebru Kizilkilic, Hakan Ozdogu, Mahmut Yeral

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DIFFERENTIAL EXPRESSION OF 16 APOPTOSIS RELATED GENES IN VITAMIN D-INDUCED DIFFERENTIATION OF HL-60 CELLS.

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COLONY FORMING ASSAY IN PATIENTS WITH AML M7

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Supportive care by granulocyte colony-stimulating factor in hypoplastic acute myelogenous leukemia

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EMA AS SALVAGE REGIMEN IN PATIENTS WITH REFRACTORY/RELAPSE AML: A SINGLE CENTER EXPERIENCE

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1Ramin Yaghobi, 2Mehdi Roshan Nia Jahromi, 3Mani Ramzi, 4Narges Rezaee, 5Vida Moaied
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1Sibel Kabukçu Hacıoğlu, 1Ismail Sari, 2Sami Kar, 3Sinemis Yüksel, 4Nilay Şen
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ARG: A POTENTIAL BIOMARKER FOR DLBCL STAGING
1Mansoor Salehi, 2Zahra Kabiri, 3Mohammad Modaresi
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LYMPHOMA EXPERIENCE OF LAKES DISTRICT FROM SÜLEYMAN DEMIREL UNIVERSITY SCHOOL OF MEDICINE
1Güçhan Alanoğlu, 2Bülent Kara, 3Sema Sezgin Gökşu, 4Nilgün Kapucuoğlu, 5Hasan Senol Coşkun
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UTILITY OF PERIPHERAL BLOOD FLOW CYTOMETRY TO INVESTIGATE THE PERIPHERIZATION OF B-CELL MALIGNANT LYMPHOMAS
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EVALUATION THE RESPONSE RATE OF IEV REGIMEN AS SALVAGE THERAPY FOR RELAPSED / REFRACTORY NON-HODGKIN’S LYMPHOMA PATIENTS
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1Evrim Kuş, 2Cengiz Erçin, 3Funda Çorapçıoğlu
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OPPORTUNISTIC INFECTIONS IN CASES WITH HAIRY CELL LEUKEMIA
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Lymhoma Experience of Lakes District from Süleyman Demirel University School of Medicine
1Güçhan Alanoğlu, 2Bülent Kara, 3Sema Sezgin Gökşu, 4Nilgün Kapucuoğlu, 5Hasan Senol Coşkun
1Süleyman Demirel University School of Medicine Dept. of Hematology, Isparta, Turkey 2Süleyman Demirel University School of Medicine Department of Internal Medicine, Isparta, Turkey 3Süleyman Demirel University School of Medicine Department of Pathology, Isparta, Turkey 4Süleyman Demirel University School of Medicine Department of Medical Oncology, Isparta, Turkey

Utility of Peripheral Blood Flow Cytometry to Investigate the Peripherization of B-Cell Malignant Lymphomas
Olga Meltem Akay, Eren Gunduz, Hava U. Teke, Gulcihan Demirel, Zafer Gulbas
Eskisehir Osmangazi University Medical School Hematology Department, Isparta, Turkey

Evaluation the Response Rate of IEV Regimen as Salvage Therapy for Relapsed / Refractory Non-Hodgkin’s Lymphoma Patients
1Mohammad Ali Mashhadi, 2Kourosh Shahraki, 3Adineh Pour
1Ali Ebne Abitaleb Hospital, Zahedan, Iran, 2Zahedan Medical University, Zahedan, Iran 3Resident of Internal Medicine in Zahedan Medical University, Zahedan, Iran

Opportunistic Infections in Cases with Hairy Cell Leukemia
Mahmut Yeral, Hakan Ozdogu, Can Boga
Baskent University Faculty of Medicine, Department of Hematology, Ankara, Turkey

Autoimmune Hemolytic Anemia After Cladribine Therapy for Hairy Cell Leukemia
Mustafa Yenerel, Esra Hatipoğlu, Abdullah Özkök, Tanju Atamer
Istanbul University, Istanbul Faculty of Medicine, Department of Internal Medicine, Division of Hematology, Istanbul, Turkey

Chronic Lymphocytic Leukemia (125)
MULTIPLE MYELOMA

Ref. No: 10 Abstract No: 62
A PROTEAZOM INHIBITOR IN THE TREATMENT OF MULTIPLE MYELOMA: BORTEZOMIB
Ozlem Sahin Balcik, Simten Dagdas, Murat Albayrak, Osman Yokusu, Funda Ceran, Servet Erbasi, Gulsum Ozet
Ankara Numune Educational and Research Hospital Hematology Department, Ankara, Turkey

Ref. No: 11 Abstract No: 63
GENETIC ABNORMALITIES IN MULTIPLE MYELOMA, THEIR PREVALENCE AND RELATION WITH OTHER RISK FACTORS
Ozlem Sahin Balcik, Murat Albayrak, Simten Dagdas, Funda Ceran, Osman Yokusu, Gulsum Ozet
Ankara Numune Educational and Research Hospital Hematology Department, Ankara, Turkey

Ref. No: 17 Abstract No: 64
TC-99M MIBI OR F-18 FDG IMAGING?: A COMPARATIVE STUDY FOR EVALUATING PATIENTS WITH MULTIPLE MYELOMA
Ilknur Ak, Inci Uslu, Zafer Gulbas
Esiksehir Osmangazi University Medical Faculty Department of Nuclear Medicine, Eskisehir, Turkey
Esiksehir Osmangazi University Medical Faculty Department of Haematology, Eskisehir, Turkey

Ref. No: 24 Abstract No: 65
BORTEZOMIB AND DEXAMETHASONE INDUCED TUMOR LYSIS SYNDROME IN A CASE OF PLASMA CELL LEUKEMIA
Gul Ilhan, Nesilhan And, Sema Karaku
Baskent University, Hematology Department, Ankara, Turkey

Ref. No: 38 Abstract No: 66
SUCCESSFUL TREATMENT OF EARLY RELAPSE OF OCULAR MYELOMA WITH BORTEZOMIB AND STEROID AFTER AUTOLOGOUS STEM CELL TRANSPLANTATION
Irfan Yavasoglu, Tolga Kocaturk, Gurhan Kadiyoklu, Volkan Dayanir, Yelda Dayanir, Zahit Bolaman
Adnan Menderes University, Medical Faculty, Hematology, Aydin, Turkey
Adnan Menderes University, Medical Faculty, Ophthalmology, Aydin, Turkey
Adnan Menderes University, Medical Faculty, Radiology, Aydin, Turkey

Ref. No: 44 Abstract No: 67
COMBINED THERAPY WITH BORTEZOMIB AND DEXAMETHASONE IN PATIENTS WITH RELAPSING MULTIPLE MYELOMA
Oktay Bilgir, Ferda Bilgir, Mehmet Calan, Pinar Oner, Murat Akyol, Elif Tuna
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Buca State Hospital, Internal Diseases Clinic, Izmir, Turkey

Ref. No: 66 Abstract No: 68
BORTEZOMIB EFFICIENCY IN MULTIPLE MYELOMA
Ebru Kizilkilic, Can Boga, Hakan Ozdogu, Mahmut Yeral
Baskent University Faculty of Medicine Department of Hematology, Ankara, Turkey

Ref. No: 67 Abstract No: 69
RETROSPECTIVE ANALYSIS FOR DEMOGRAPHIC FEATURES OF MULTIPLE MYELOMA PATIENTS, AKDENIZ UNIVERSITY EXPERIENCE
Mete Akin, Ilknur Nizam, Songul Akcan, Feyzi Bostan, Ihsan Karadogan, Aysen Timuragaoglu, Levent Undar
Akdeniz University Dept. of Hematology, Antalya, Turkey

Ref. No: 68 Abstract No: 70
THE EFFECTS OF PLASMA EXCHANGE ON COAGULATION PARAMETERS, AND PLATELET FUNCTIONS IN PATIENTS WITH MULTIPLE MYELOMA
Ali Sahin, Ali Uenal, Fatih Kurnaz, Leylagul Kaynar, Mehmet Oztekin, Musa Solmaz, Fevzi Altuntas, Bulent Eser, Mustafa Cetin
Erciyes University, Medical Faculty, Hematology Department, Kayseri, Turkey

Ref. No: 69 Abstract No: 71
RETROSPECTIVE ANALYSIS FOR CLINICAL AND LABORATORY FINDINGS OF MULTIPLE MYELOMA PATIENTS, AKDENIZ UNIVERSITY EXPERIENCE
Mete Akin, Ilknur Nizam, Songul Akcan, Feyzi Bostan, Ihsan Karadogan, Aysen Timuragaoglu, Levent Undar
Akdeniz University Dept. of Hematology, Antalya, Turkey

Ref. No: 73 Abstract No: 72
MICROSATELLITE INSTABILITY IN PATIENTS WITH MULTIPLE MYELOMA
Aysen Timuragaoglu, Evren Kiriş, Sema Demircan, Seray Dizlek, Guçhan Alanoglu, Nilay Uysalgil
Akdeniz University, School Of Medicine, Antalya, Turkey
Silleyman Demirel University, School Of Medicine, Isparta, Turkey

Ref. No: 75 Abstract No: 73
AN UNUSUAL PRESENTATION OF MULTIPLE MYELOMA: PLASMACYTIC ASCITES COMPLICATED BY DUODENAL INVOLVEMENT
Q meltem Akay, Baris Cansu, F Mustafa Akgalin, Emre Entok, Zafer Gulbas
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Osmangazi University Faculty of Medicine, Department of Pathology, Eskisehir, Turkey
Osmangazi University Faculty of Medicine, Department of Nuclear Medicine, Eskisehir, Turkey

Ref. No: 79 Abstract No: 74
TIME INTERVALS PRECEEDING AUTOLOGOUS STEM CELL TRANSPLANTATION (SCT) IN MULTIPLE MYELOMA PATIENTS: A SINGLE CENTER INTENT TO TRANSPLANT ANALYSIS
Mutlu Arat, Merih Kizil Cakar, Ender Soydan, Pervin Topcuoglu, Aynur Ugur Bilgin, Sule Mine Bakanay, Erol Ayyildiz, Onder Arslan, Muhit Ozcan, Gunhan Gurman, Meral Bekac, Osman Ilhan
Ankara University, School of Medicine, Department of Hematology, Ankara, Turkey

Ref. No: 81 Abstract No: 75
MULTIPLE MYELOMA WITH MASSIVE ASCITES: A CASE REPORT
Nurhial Turgut, Inci Alacacioglu, Ozden Piskin, Selda Ceneli, Guner Hayri Ozsan, Fatih Demirkiran, Mehmet Ali Ozcan, Bulent Undar
Dokuz Eylul University Faculty of Medicine Department of Hematology, Izmir, Turkey

Ref. No: 87 Abstract No: 76
RETROSPECTIVE ANALYSIS FOR CLINICAL FEATURES OF MULTIPLE MYELOMA PATIENTS, AKDENIZ UNIVERSITY EXPERIENCE
Mete Akin, Ilknur Nizam, Songul Akcan, Feyzi Bostan, Ihsan Karadogan, Aysen Timuragaoglu, Levent Undar
Akdeniz University Dept. of Hematology, Antalya, Turkey
TREATMENT OF RELAPSED/REFRACTORY MULTIPLE MYELOMA WITH THALIDOMIDE: A RETROSPECTIVE EVALUATION IN A CENTER
1Bahriye Payzin, 2Gülbin Seyman Çetinkaya
1İzmir Atatürk Research Training Hospital, Department of Hematology, Izmir, Turkey 2İzmir Atatürk Research Training Hospital, Department of Internal Medicine, Izmir, Turkey

Ref. No: 103 Abstract No: 77

MULTIPLE MYELOMA: RETROSPECTIVE ANALYSIS OF 35 PATIENTS
Gülten Sop, Füsun Özdemirkıran, Tuğba Gümüş, Şermin Coşan
İzmir Training And Research Hospital, Izmir, Turkey

Ref. No: 113 Abstract No: 78

ORAL MELPHALAN AND PREDNISONE PLUS THALIDOMIDE COMPARED WITH HIGH-DOSE THERAPY FOLLOWED BY AUTOLOGOUS PERIPHERAL-BLOOD STEM-CELL TRANSPLANTATION IN PATIENTS WITH MULTIPLE MYELOMA
Hakan Özdogu, Can Boga, Ebru Kızılkılık, Mahmut Yeral
Baskent University Faculty Of Medicine, Department of Hematology, Ankara, Turkey

Ref. No: 132 Abstract No: 79

CLINICAL AND BIOCHEMICAL FEATURES FOR MONITORING MULTIPLE MYELOMA: A RETROSPECTIVE ANALYSIS FROM “DENIZLI LEUKEMIA-LYMPHOMA-MYELOMA STUDY GROUP” (DLMSSG)
1Sibel Kabukçu Hacıoğlu, 1İsmail Sari, 2Sami Kartı, 3Nilay Sen, 4Belda Dursun, 1Ali Keskin
1Pamukkale University, Faculty of Medicine, Department of Hematology, Denizli, Turkey 2Denizli Education and Research Hospital, Hematology Unit, Denizli, Turkey 3Pamukkale University, Faculty of Medicine, Department of Pathology, Denizli, Turkey 4Pamukkale University, Faculty of Medicine, Department of Nephrology, Denizli, Turkey

Ref. No: 98 Abstract No: 80

ENDOTHELIAL CELL KINETICS IN PLASMA CELL LEUKEMIA
1İlkınur Kozanoglu, 2Hakan Özdogu, 2Can Boga, 3Erkan Maytalman, 4Oktay Sözer
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OSTEONECROSIS OF THE JAW IN PATIENTS WITH MULTIPLE MYELOMA TREATED WITH ZOLEDRONIC ACID
Selât Çetiner, Gülşan Türköz Sucak, Şahika Zeynep Ak, Benay Kocakahyaoglu, Selvi Kahraman, Mehmet Araç, Mustafa Çetiner, Erta Delilloçoğlu, Rauf Haznedar
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4Baskent University Adana Hospital Hematology Research Laboratory, Adana, Turkey

Ref. No: 115 Abstract No: 87

DOES BRASSICA RAPA (A PLANT FROM FAMILY BRASSICACEAE) SOLUTION INDUCE FLUID RETENTION IN CML PATIENTS WHO RECEIVED IMATINIB?
Suheyl Asma, Can Boga, Hakan Ozdogu
Baskent University Faculty of Medicine, Department of Hematology, Ankara, Turkey

Ref. No: 117 Abstract No: 88

TREATMENT ALTERNATIVES IN YOUNG-POOR RISK CML PATIENTS
Suheyl Asma, Can Boga, Hakan Ozdogu, Ebru Kızılkılık
Baskent University Faculty of Medicine, Department of Hematology, Ankara, Turkey

MYELODYSPLASTIC SYNDROMES (134-135)

Ref. No: 3 Abstract No: 82

DATA FROM THE REGISTRY OF THE PATIENTS WITH MYELODYSPLASTIC SYNDROME FROM CLINIC OF HEMATOLOGY, FUNDENI CLINICAL INSTITUTE, BUCHAREST, ROMANIA. I. EPIDEMIOLOGICAL GENERAL DATA
Gologan Radu, Georgescu Daniela
Clinic of Hematology, Fundeni Clinical Institute, Bucharest, Romania

Ref. No: 108 Abstract No: 83

TREATMENT OF HIGH-RISK MYELODYSPLASTIC SYNDROME WITH DEMETHYLATING AGENTS
Banu Diri, Can Boga, Hakan Ozdogu, Mutlu Kasar
Baskent University Faculty Of Medicine, Department Of Hematology, Adana, Turkey

Ref. No: 134 Abstract No: 84

FLOW CYTOMETRIC ANALYSIS OF PERIPHERAL BLOOD IN DIAGNOSIS OF MYELODYSPLASTIC SYNDROMES
Eren Gündüz, Olga Meltem Akay, Hava Üskûdar Teke, Gülçihan Demirel, Zafer Gülbaş
Osmangazi University, Eskişehir, Turkey

MYELOPROLIFERATIVE DISORDERS (135)

Ref. No: 97 Abstract No: 85

CIRCULATING CD34 CELLS IN MYELOFIBROSIS
1İlkınur Kozanoglu, 2Hakan Özdogu, 3Can Boğa, 4Oktay Sözer
1Baskent University Medical Faculty Physiology Department, Adana, Turkey 2Baskent University Medical Faculty Hematology Department, Adana, Turkey 3Baskent University Adana Hospital Hematology Research Laboratory, Adana, Turkey

Ref. No: 107 Abstract No: 86

NORMALIZATION OF PLATELET COUNT DURING PREGNANCY IN A PATIENT WITH ESSENTIAL THROMBOCYTHEMIA
Can Boğa, Hakan Özdogu
Baskent University Faculty of Medicine, Department of Hematology, Ankara, Turkey

CHRONIC MYELOID LEUKEMIA (135-136)

Ref. No: 115 Abstract No: 87

DOES BRASSICA RAPA (A PLANT FROM FAMILY BRASSICACEAE) SOLUTION INDUCE FLUID RETENTION IN CML PATIENTS WHO RECEIVED IMATINIB?
Suheyl Asma, Can Boga, Hakan Ozdogu
Baskent University Faculty of Medicine, Department of Hematology, Ankara, Turkey

Ref. No: 117 Abstract No: 88

TREATMENT ALTERNATIVES IN YOUNG-POOR RISK CML PATIENTS
Suheyl Asma, Can Boga, Hakan Ozdogu, Ebru Kızılkılık
Baskent University Faculty of Medicine, Department of Hematology, Ankara, Turkey
PERIPHERAL POLYNEUROPATHY ASSOCIATED WITH IMATINIB TREATMENT
Can Boga, Hakan Ozdogu
Baskent University Faculty of Medicine, Department of Hematology, Ankara, Turkey

PALLIATIVE CARE – SUPPORTIVE THERAPY (136-137)

FEVERED NEUTROPENIC EPISODES IN ACUTE LEUKEMIA PATIENTS: EXPERIENCE OF BAŞKENT UNIVERSITY HOSPITAL
1Neslihan Andıc, 1Sema Karakuş, 1Gül İlhan, 2Funda Timurkaynak, 2Hande Aslan
1Baskent University, Faculty of Medicine, Department of Internal Medicine, Hematology Division, Adana, Turkey
2Baskent University, Faculty of Medicine, Department of Infectious Diseases, Ankara, Turkey

STEM CELL TRANSPANTATION (137)

TESTS, HISTORICAL EFFORTS AND IMMUNE RECONSTITUTION IN CORD BLOOD STEM CELLS TRANSPLANTATION
Shaban Alizadeh, Ali Abedi, Shahab Bohlooli
Arums, Iran

AUTOLOGOUS HEMATOPOIETIC STEM CELL TRANSPLANTATION FOR LYMPHOMA AND MYELOMA AT YEDITEPE UNIVERSITY HOSPITAL
Sabih Yuce, Gulcin Kalayci, Sema Aktas, Didem Aydin, Basak Oyan, Yener Koc
Yeditepe University Stem Cell Transplant Unit, Istanbul, Turkey

RELATIONSHIP BETWEEN INSULIN RESISTANCE AND SOME COAGULATION AND FIBRINOLYTIC PARAMETERS IN SUBJECTS WITH METABOLIC SYNDROME
1Nashwa Abou Samra, 1Amany Ragab, 2Asmaa Higazy, 2Omayma Saleh
1Departments Of Clinical Pathology Faculty Of Medicine, Mansoura University, Egypt 2Internal Medicine, Faculty Of Medicine, Mansoura University, Egypt
ACUTE LYMPHOBLASTIC LEUKEMIA

Ref: No: 8 Abstract No: 1

THALASSEMA MINOR AS ONE OF RISK FACTORS FOR CHILDHOOD LEUKEMIA

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Leukemia is a highly frequent cancer with an unknown mechanism of pathogenesis in the most of the cases. This study was performed to determine the association of some probable life threatening factors with childhood leukemia. 153 bone marrow aspiration (BMA) confirmed leukemic children (age<18 years) and 153 age-sex matched healthy controls from Southern Iran were asked about probable risk factors for leukemia. 88 of the patients who were in remission phase were studied for thalassemia trait according to complete blood count and hemoglobin electrophoresis. Significant differences were observed between patients and controls in birth weight (P<0.003), maternal age during pregnancy (P<0.001), life closed to powerful electrical stations (P<0.002), Pregnant mother or child passive smoking (P<0.001), father injuring with chemical weapons (P<0.005) and family history of leukemia (P<0.02) while encountering of mother and child with X-ray, history of other malignancies in child and history of abortion were not statistically different. Frequency of thalassemia minor in the patients was higher than normal population (23.9% vs. 7.5%, P<0.001, odds ratio=3.87, 1.87<OR.

Ref: No: 13 Abstract No: 2

LATE EXTRAMEDULLARY RELAPSES OF ACUTE LEUKEMIA

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Childrens Hospital of University Mainz, Department of Pediatric Surgery, Germany

Introduction: We report about two girls with isolated ovarian extramedullary relapse of acute lymphoblastic leukemia. We present the case histories and we will discuss the role of therapy (chemotherapy and radiati) and of surgery in these cases. Case 1: This girl had the primary diagnosis of a Pre-B-All (acute lymphoblastic leukemia, immunologically typed as “low risk”). Treatment was done according to the CoALL82-protocol, which is one of the two usually used treatment protocols for ALL in Germany. At the age of 7 and 9 years bone marrow relapses occurred. They were retreated by even more aggressive chemotherapy till the age of 11 years. After a while she presented with a huge abdominal mass, the open biopsy showed lymphoblastic cells in the ovary, chemotherapy and low-dose radiatio. Salpingo-oophorectomy was done. 21 years after the first diagnosis of ALL, the young women is in complete continuous hematologic remission. Case 2: 14 years old girl, with Pre-B-All, typ “low risk” was treated according to the treatment protocol CoALL06-97. She was in hematologic remission for 18 months. Then she had abdominal pain. Ultrasound and CT showed masses originating from both ovaries (5 and 8 cm). In the laparoscopic biopsies from one ovary and a lymph node we found ovarian relapse of ALL, in the bone marrow puncture, which was done during the operation, were no signs of malignancy. The treatment was following BFM protocol for ALL-recurrence. The tumor masses regressed and after 6 months control laparoscopy was done. At the age of 18 years she has ophthalmological problems and now we diagnosed a isolated CNS-relapse of ALL. Discussion: In the second case additional radiatio as a potential treatment modality was avoided and fertility probably preserved. Laparoscopy is able to explore abdominal tumors, do staging procedures, and it can give a very good visualisation of intraabdominal tumors. In selected cases biopsies can be done; the indication can intraoperatively be discussed interdisciplinary. This way of exploration of abdominal gives safer results of histologic specimen compared with percutaneous biopsy and it is less invasive compared with open biopsy.

Ref: No: 83 Abstract No: 3

RESULTS OF HYPERFRACTIONATED CYCLOPHOSPHAMIDE, VINCristINE, DOxorubicin, and DEXAMETHASONE (HYPER-CvAD) IN ADULT ACUTE LYMPHOBLASTIC LEUKEMIA: SINGLE CENTER EXPERIENCE

Inci Alacacioglu, Nurhilal Tunurt, Fatih Demirkan, Mehmet Ali Ozcan, Ozden Pişkin, Güner Hayri Ozsan, Selda Ceneli, Bülent Ündar

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Objective: The aim of the study was evaluation of the effect of hyperfractionated cyclophosphamide, vincristine, doxorubicin, and dexamethasone (Hyper-CvAD) regimen on remission-induction and overall survival (OS) in acute lymphoblastic leukemia (ALL) patients. Methods: Twenty patients with ALL who were treated with Hyper-CvAD between January 2001 and May 2006 in our clinic were evaluated retrospectively. The median age of the patients at diagnosis was 32±13. 1 with a 3/1 M/F ratio. 60% (12 patients) was diagnosed as precursor B cell, 5% (1 patient) as mature B cell, 35% (7 patients), as T-cell ALL. The overall incidence of Philadelphia chromosome (Ph) positive ALL was 15%. Results: The mean follow-up time of all patients was 14. 1 months (3-36 months). A complete response (CR) was achieved in 75% of the patients. Duration of CR was 12±7. 8 months in these patients. The induction mortality rate was 5% (1/20). The OS was 14±1. 8 months with a 20% 2-year survival rate. The patients with good performance status (ECOG 1-2) lived longer (21±5. 57months x 8±1. 5months, p=0. 03). Conclusion: Short term follow-up results with hyper-CvAD regimen does not seem favorable according to our experience.

Ref: No: 86 Abstract No: 4

METHOTREXATE INDUCED NEUROTOXICITY DURING HYPER CVAD TREATMENT IN A PATIENT WITH ALL

Düzgün Ozatli, Nil Güler, Nevzat Selim, Mehmet Turgut

Ondokuz Mayis University, Samsun, Turkey

Methotrexate induced neurotoxicity during Hyper CVAD treatment in a patient with ALL in the group of patient with ALL. Düzgün Özatli, Nil Güler, Nevzat Selim, Mehmet Turgut Ondokuz Mayis
University, Medical School, Department of Hematology, Samsun, Turkey Methotrexate related neurotoxicity is well documented among children ALL patients. It is frequently presented as seizures. Most authors believe that neurotoxicity induced with methotrexate may be due to complex or multifactorial mechanisms. Here in, we will present young adult man complicated his treatment with neuropathy occurring during methotrexate infusion. Twenty years old man diagnosed with T cell acute lymphoblastic leukemia (ALL). Hyper CVAD A chemotherapy protocol treatment was given. His bone marrow aspiration was compatible with remission after chemotherapy protocol treatment. He was hospitalised again for Hyper CVAD B chemotherapy protocol treatment. The symptoms that compatible with periferic facial paralysis was occurred during methotrexate loading dose infusion. Treatment was stopped. Ca,Mg,K levels were normal. Symptoms regressed after stopping the methotrexate treatment. After one hour, methotrexate infusion treatment was given the patient again. Same symptoms were occurred. The treatment was stopped again. A neurological consultation was made. His cranial CT and MR were normal. Neurological examination was compatible with periferic facial paralysis. It resolved with steroids within days. Our consideration is this situation was result of the methotrexate neurotoxicity. We found this case report as valuable. Because the methotrexate induced neurotoxicity is frequently presented as seizures.

METHOTREXATE INDUCED NEUROTOXICITY DURING HYPER CVAD TREATMENT IN A PATIENT WITH ALL
Düzgün Özathi, Nil Güler, Nevzat Selim, Nazir Yayla
Ondokuz Mayıs University, Samsun, Turkey
Methotrexate induced neurotoxicity during Hyper CVAD treatment in a patient with ALL Düzgün Özathi, Nil Güler, Nevzat Selim, Nazir Yayla Ondokuz Mayıs University, Medical School, Department of Hematology, Samsun, Turkey Methotrexate related neurotoxicity is well documented among children acute lymphoblastic leukemia (ALL) patients. It is frequently presented as seizures. Most authors believe that neurotoxicity induced with methotrexate may be due to complex or multifactorial mechanisms. Here in, we will present young adult man complicated his treatment with neuropathy occurring during methotrexate infusion. Twenty years old man diagnosed T cell ALL. Hyper CVAD chemotherapy treatment was given. After A arm of this protocol treatment his bone marrow aspiration was compatible with remission. During Hyper CVAD B arm chemotherapy treatment, the symptoms that compatible with periferic facial paralysis was occurred during methotrexate loading dose infusion. Treatment was stopped. All blood electrolyte levels were normal limits. Symptoms regressed after stopping the methotrexate treatment. After one hour, methotrexate infusion treatment was given the patient again. Same symptoms were occurred. The treatment was stopped again. A neurological consultation was made. Neurological examination was compatible with periferic facial paralysis. His cranial CT and MR were normal. Steroid treatment was started and the symptoms were resolved within days. Our consideration is that the situation was result of the methotrexate neurotoxicity. This is the first case presented with peripheral facial paralysis due to methotrexate neurotoxicity.

RESEARCHING THE EFFECT OF ANTHRACYCLINE CARDIOTOXICITY TO SYSTOLIC AND DIASTOLIC FUNCTIONS OF HEART WITH ECHOCARDIOGRAPHY
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Objective: The objective of this study was to examine the effects of anthracycline treatment to diastolic and systolic functions of heart by echocardiographic studies. Methods: Between March 2002 and February 2003, 25 children in the care of Bakırköy Maternity and Children Hospital, receiving chemotherapy were examined. Studies were stratified according to cumulative anthracycline dose into five groups. Diastolic and systolic functions of heart were determined for each patient before and during treatment by Doppler echocardiography. Results: The study included 25 patients. There were 13 girls and 12 boys. 15 children (7 boys and 8 girls) were taken to control group. Body surface area and ages of patient and control groups were similar. Blood pressure, left ventricular diastolic and systolic diameter of pretreatment group and control group were not similar. During treatment increase at heart rate, systolic blood pressure, increase a light diastolic filling velocity and decrease at E/A ratio of patient and control groups were not similar. Conclusions: The determination of the changes at the diastolic functions is important for the diagnosis early heart disease. Further research is needed to determine these datas.

DETERMINING EARLY ANTHRACYCLINE TOXICITY WITH ECHOCARDIOGRAPHIC STUDIES AND CARDIAC TROPONIN I
Kazım Öztarhan, Meliha Aslan, Belgin Aktaş, Gönül Aydoğan, Zafer Salçoğlu, Ferhan Akcioğlu
Bakırköy Maternity and Children Hospital, İstanbul, Turkey
Abstract: Childhood acute leukemia is the most common malignancy diagnosed in children (%32). In our country after leukemia the second common malignancy is lymphome(%25. 3) and the third is central nervous system tumors(%10. 6). By the last 30 years with the development in the diagnosis methods especially at lymphoblastic leukemia the long time survival percentage reached up to %70-80. Anthracycline drugs are given as part of chemotherapy for a wide range of malignant diseases and have been particularly valuable for the successful treatment of many childhood malignancies. The most commonly used types are doxorubicin and daunorubicin. Because of the high antitumoral effect, doxorubicin is used at the treatment of either solid tumors or hematologic malignancies. Daunorubicin is used effectively at the treatment of acute lymphoblastic and myeloblastic leukemia for both children and adults. More than 20 years the anthracycline cardiotoxicity is associated with cumulative doses. This threats the patients cardiac functions and limits the use of these drugs. Clinically the cardiotoxicity related with dose can be seen weeks, months or years after treatment ends. To determine the cardiotoxic effects of these agents some of the recent used methods are: following cardiac enzymes, electrocardiography, echocardiography, radionucleid angiography and transvenoz cardiac biopsy. Results: 158 echocardiographic and cardiac troponin studies were performed. Studies
were stratified according to cumulative anthracycline dose into nine groups. Diastolic and systolic functions of heart were determined for each patient before and during treatment by Doppler echocardiography and blood cardiac troponin. The importance of echocardiography and cardiac troponin by earlier diagnosing of cardiotoxicity is determined. Cardiac failure is occurred at 8 patient during chemotherapy. Although antikongestive treatment 2 patients were died. Both of them were ALL-L1 diagnosed girls. Conclusions: 1- Recent studies shows that before systolic functions diastolic functions are damaged. In our study we had the similar results to these studies. In our study peak atrial phase velocity(Av) was increased and E/A ratio was decreased. We determined the cut-off values of peak atrial phase velocity(Av) and E/A ratio. According to this Av>0.6180m/sn and E/A<1 are the cut-off values of diastolic function damages. (Fig. 1-2) 2. Our study shows that cardiac troponin doesn’t determines the myocard damage related with cumulative anthracycline dose but increases after the last term heart insufficiency begins. (Fig. 3) According to this, determining early cardiac damage related with anthracyclines cardiac troponin doesn’t have a specific role. 3. At early diagnosis of anthracycline cardiotoxicity echocardiographic studies have the most important role. Further research is needed for the cardiac troponins.

![Fig 1. Changes in E/A ratio with increasing cumulative anthracycline dose.](image)

Ref: No: 126 Abstract No: 9

ALL CASE WITH MULTIPLE SOLID LESIONS IN LIVER

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Introduction: Hepatomegaly is common in newly diagnosed and relapsed patients with acute lymphoblastic leukemia(ALL). Usually, it is related to diffuse enlargement secondary to infiltration by leukemic lymphoblasts. Involvement as solid mass lesions is rare. Since multiple solid lesions in liver cases are not common, we have found this case suitable to declare. Case: 22 year old woman was diagnosed B-ALL a year ago. We pointed bone marrow relapse and planned to apply allogeneic hematopoietic stem cell transplantation. Multiple solid lesions in liver were detected by ultrasonography just before treatment. Biopsy showed ALL infiltration. In addition hematological relapse was observed. FLAG treatment protocol including fludarabine and ARA-C was given to the patient. In bone marrow examination after treatment, we established remission. And there were no lesions in liver by ultrasonography.

Ref: No: 128 Abstract No: 9

ACUTE LYMPHOBLASTIC LEUKEMIA

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Abstract: we performed this study during 10 years, at the mentioned hospital From 1987 until 1996. During this time studying performed on 76 leukemic Children at that Hospital. From 76 cases acute leukemias, 7 cases Were Acute Myeloid leukemias and 69 cases were ALL. From 69 cases Acute Lymphoblastic leukemias, 29 cases were ALL- L1, and 34 cases Were ALL-L2 and 6 cases were only ALL- L3. The blast cells in ALL-L3 Were large /uniform, and nucleus shape was round to oval,homogenous, And had also one or more,vesicular, often prominent nucleoli and its Cytoplasm, abundant,deeply basophilic. Aims: Studing on ALL burkit Type and its causes. Methods: Examinations peripheral blood and B. M. and cytochemical methods, besides Cytochemical methods, immunologic procedures have proved very useful for Classifying the acute leukemias and especially for the differentiation of ALL And its subgroups. Results: This type of ALL is rare and usually are B- ALL Conclusions: This type of ALL was seen in children who had 9 to 11 or 12 years old and it sounds that they was born from older mothers Or women, because, we observed that the their mothers had over 40 years old. Email: hem1331@yahoo.com

Ref: No: 128 Abstract No: 9

FREQUENCY OF CANCERS IN CHILDREN UNDER 14 YEARS OLD IN ALI EBNE ABITALEB HOSPITAL IN ZAHEDAN 2003-2006

Mohammad Ali Mashhadi, Kourosh Shahraki, Eghbal Shizraei, Parnoush Tajbakhsh, Rahime Khademi, Alireza Rezvani, Neda Shahraki

1Ali Ebne Abitaleb Hospital Zahedan, Iran, 2Zahedan Medical University, Zahedan, Iran

Background: The aim of this study is to present the frequency of cancers in children who were treated in Ali Ebne Abitaleb Hospital of the Zahedan in the period of 2003-2006. Patients & Methods: This retrospective survey covered consecutively diagnosed and treated patients admitted to Ali Ebne Abitaleb Hospital in Zahedan. The research protocol was discussed extensively, so the data to be collected and the criteria for their evaluation were clearly pre-defined. We analyzed 714 patients diagnosed between 2003-2006 with various cancers. 147 cases under 14 years old was selected. From these analyses, the general and specific frequency by age and by sex were obtained for the different group of neoplasms. Also, the frequency of the stage of the disease that had been diagnosed in cases of children with solid tumors was obtained. Survival analyses were carried out using the SPSS and Kaplan-Meier Method, according to gender, age, vital status and stage. Results: A total 147 cases of children with cancer were diagnosed, with the male/female ratio at 1. 7/1. Leukemia had the highest frequency with 94 cases (63%) and, of these cases, Acute Lymphoblastic Leukemia (ALL) was the most prevalent with 77 cases (82%). Thereafter, in descending order of frequency, were Lymphoma with 27 cases(18%); HD=9 & NHL=18, Bone

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ACUTE MYELOBLASTIC LEUKEMIA

Ref: No: 43 Abstract No: 11

ABBERANT ANTIGEN EXPRESSION IN 236 PATIENTS WITH ACUTE MYELOID LEUKEMIA BY MULTI-COLOR FLOW CYTOMETRY

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1. Ankara Numune Education And Research Hospital, Ankara, Turkey 2. Ufuk University, Ankara, Turkey 3. Karaman University, Zonguldak, Turkey

Recently immunophenotyping is spredely used in the diagnosis and classification of leukemia. To evaluate the immunophenotype and abberant antigen expression of acute myeloid leukaemi (AML), multiparameter flow cytometry and CD45/SSC gating were used to analyse the surface and cytoplasmic antigen expression in 236 new diagnosed AML patients in Ankara Numune Education and Research Hospital between 2003-2005. The results were compared with FAB classification to help define the best use and role of multiparameter flow cytometry in the diagnosis and proper classification of AML. The distribution of the patients according to FAB classification was as follows: 30 (13%) patients were M0, 60 (26%) were M1, 80 (34%) were M2, 21 (9%) were M3, 27 (12%) were M4, 11 (5%) were M5 and 7 (3%) were M6. CD34, CD117, CD13, CD33, CD12, CD11b, CD115, CD64, cMPO, cTdT, Anti-CD1HAD, CD2, CD7, CD19, CD56 expressions were analysed in all of the patients’ marrow aspiration materials or blood samples by flow cytometry. CD56 was the most commonly expressed abberant antigen (19%), followed by CD7 (12%), CD7+CD56 (8%), CD19 (6%), CD2 (4%), CD7+CD19+ (3%). Some immunophenotypes correlated with FAB type, including increased frequency of CD2 and CD56 in M3; increased frequency of CD19 in M2, and CD7+CD56 in M0.

Ref: No: 59 Abstract No: 12

RETROSPECTIVE ANALYSIS OF ACUTE PROMYELOCYTIC LEUKEMIA PATIENTS IN OUR CENTER

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Acute promyelocytic leukemia comprises approximately 10-15% of acute myeloid leukemia. The disease is more often at younger age in contrast to the other acute myeloid leukemia subtypes and the most significant characteristic is that there is an increased tendency to hemorrhage. By the late 1980’s the disease had been treated like the other acute myeloid leukemia subtypes and complete remission was reported in about 60-65% of the cases. In our study, we aimed to investigate the data of the patients with acute promyelocytic leukemia retrospectively between 2000-2006 and to determine our own findings and compare the compatibility of them with literature. Modify AIDA protocol was administrated to all the patients. The median age of the group was 42 and it was younger than that of acute myeloid leukemia. There was no difference between the gender. The patients average application time to hospital was 11 days after the symptoms occurred. The medium white cell count was 23, 180/mm3 which was 2000/mm3 much higher than the literature. %89 of patients at admission had symptoms and signs (ecchymosis, hematuria, epistaxis) of bleeding tendency. However, we demonstrated coagulopathy at %51 of patients by laboratory. The coagulopathy clinic and laboratory symptoms were concordant with the literature. The complete remission rate of our patients was %70.8. Four patients (%17) died due to early hemorrhagic complications. The complete remission was lower and early death rate was higher when compared with the literature. Retinoic acid syndrome rate was %22. Retinoic acid syndrome rate is reported to be 10-15 in literature. Even if white blood cell count on admission to hospital is reported to be frequently high, there is no precise risk factor that has been demonstrated for the development of the retinoic acid syndrome. We also have failed to find a parameter leading to predisposition to the syndrome. Consequently, there was no difference between the patient group and the literature in terms of the demographic findings such as age, gender. The development of retinoic acid syndrome and the loss of patients resulting from early death were higher, but complete remission rate was lower compared to literature. The reasons behind these were that the white blood cell count on admission to hospital were higher compared with the literature and the patients were referred to hospital late. The difference between literature and our findings might be caused by the fact that the number of our patients were lower than that of the multicenter study.

Ref: No: 62 Abstract No: 13

CD33-DIRECTED THERAPY WITH GEMTUZUMAB OZOGAMICIN IN ACUTE MYELOID LEUKEMIA: REPORT OF TWO CASES

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Baskent University Faculty of Medicine, Department of Hematology, Ankara, Turkey

Acute myeloid leukemia is a heterogeneous group of haematopoietic clonal disorders. In most cases, a combination of anthracylin and cytarabine is used during the induction phase. Unfortunately many patients relapse within a few months. CD33 is expressed on the malignant tumors with 8 cases (5%), Tumors of the Central Nervous System (CNST) with 7 cases (4%), Neuroblastoma with 4 cases (2%). The highest frequency of cancer was found in the group of 4.8 years olds that had 53 cases (36%) and in the group of 0.4 years; N=47, 31%. In the all age groups, leukemia was the most frequent. Of those cases of solid tumors for which the stage of the disease had been determined 64% were diagnosed as being stage III or IV. Conclusions: The principals cancers in the children treated in Ali Ebne Abitaleb Hospital of Zahedan were Leukemia, Lymphoma, and CNST consistent with those reported by other place. In this population Leukemia had a very high incidence and that for Germal Cell Tumor and Neuroblastoma is very low. This fact will need to be confirmed by a longer period of observation, but even now the total number of cases (particularly Leukemia) is high when compared with the data of other children Leukemia regisries which give rates for longer period and for similar or larger population.
blast cells in most cases of acute myeloid leukemia (AML) but not on normal hematopoietic pluripotent stem cells. Antibody-based therapies have focused on the membrane antigen CD33. Gemtuzumab ozogamicin consists of a humanized IgG4 anti CD33 monoclonal antibody joined to N-acetyl-g-calicheamicin dimethyl hydrazide. We report the use of single agent gemtuzumab ozogamicin (GO) in chemotherapy refractory relapse acute myeloid leukemia. A 17-years old boy was diagnosed with AML MO in May 2005. Cytogenetic analysis was normal. Induction treatment consisted of a combination of idarubicin and cytarabine. He was not response for induction therapy. He achieved partial response after fludarabine, cytarabine and G-CSF (FLAG). He had identical related donor (his aunt) and peripheral stem cell transplantation was performed in March 2006. He relapsed on day +95. The patient received donor lymphocyte infusion and high dose cytarabine. This salvage therapy failed and a bone marrow aspirate a month later showed massive infiltration of blast cells. The positive CD33 expression by the blast use of gemtuzumab ozogamicin was proposed. This treatment was given at conventional dose of 9 mg/m², given 2h infusion on days 1 and 15. On day 35 after first GO dose, control bone marrow aspirate showed massive infiltration. He died after 3 month due to sepsis. The other patient was 60 years old man. He was diagnosed with AML MO in April 2006. Cytogenetic revealed the presence of deletion 7q. He did not response induction therapy with cytarabine and mitoxantrone. The patient was given 9 mg/m² gemtuzumab ozogamicin. He died after one day with sepsis. This observation supported the thesis that GO might be useful in patients with AML before deterioration.

Ref. No: 71  Abstract No: 14

INTRACRANIAL GRANULOCYTIC SARCOMA CASE
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Ankara Numune Hospital, Ankara, Turkey

Introduction: Granulacytic sarcoma (GS) is an extra-medullary tumor with immature myeloid cells. It can be seen as a complication during Acute Myeloid Leukemia (AML), Myelodysplastic Syndrome or Myeloproliferative Diseases. Many systems could be involved such as skin, lungs, genitourinary system and breasts. In central nervous system, cerebro spinal fluid invasion could be seen but parachimal lesions are rare. Generally GS’s are divided in 4 groups: 1) Primary GS 2) GS in AML 3) GS as isolated relapse 4) GS present at the diagnoses of AML. Intracerebral GS has a poor prognosis. Diagnosis is made by performing a biopsi on the mass detected by MR or CT. Systemic treatment that contains high dose ARA-C, local treatment (intrathecal) and radiotherapy could be helpful. As intracranial GS cases are rare we have found this case suitable to declare. CASE: 35 year old man, who was diagnosed AML 2 years ago, was in remission when he came up in february 2007 with headache, unable to walk and changes in conciousness. At the admission we noted bone marrow relapse. With Cranial CT a tumor was detected in right posterior fossa which is 40x32 mm sized, surrounded with edema and compressing the fourth ventricule. Also MRI was performed. Steroids and anti-edematous treatment was given at first but there was no significant difference in the clinic of the patient. Patient couldn’t be operated or any biopsy couldn’t be taken from the lesion because of his general situation. We initiated the high dose ARA-C therapy. In the second day of the treatment patients headaches started to heal and his general health showed improvement. In the following days the patient was able to walk again. In bone marrow examination after treatment, we established remission. Significant regression was seen in control CT and MRI. Under this circumstances we diagnosed patient as GS.

Ref. No: 78  Abstract No: 15

THE DETERMINATION OF LEUKEMIC PHENOTYPE AND MINIMAL RESIDUAL DISEASE IN ACUTE MYELOID LEUKEMIA BY USING FLOW CYTOMETRY
Perin Topcuoglu, Klara Dalva, Sema Merc, Sema Ipek, Sahin, Meral Bekas, Mutlu Arat
Ankara University, School of Medicine, Ankara, Turkey

The hematological remission in acute leukemia is defined as the achievement of complete hematological recovery in peripheral blood and the a decrease of leukemic blasts below 5 %. The persistency of malignant cells below 5% is called as minimal residual disease (MRD). A number of publications have reported that the presence of MRD after the treatment(s) might predict relapse of the underlying disease. One of the convenient methods used for the detection of MRD is immunophenotyping with FCM. In this study we aimed to evaluate both the leukemic cell immunophenotype(IP) at the diagnosis and the amount of MRD at the follow-up in AML patients (using FCM events). 166 patients diagnosed as AML between January 2004 and June 2006 in our center was analyzed retrospectively. Median age was 49 years. Male/ Female was 104/62. Method: Immunophenotyping at the diagnosis was performed by using the monoclonal antibodies specific for CD45, HLA-DR, CD34, CD33, MPO, CD15(CD16), CD13, CD24, CD11b, CD117, CD64, CD10, CD2, CD19, TdT and (CD4)(CD7). The data were collected in FC500 (Beckman Coulter, France) flow cytometer and were analyzed using the RX-P software program. We usually analyzed approximately 10.000 cells at the diagnostic samples and at least 500.000 cells at the follow-up samples for the detection of MRD. The classification of MRD was evaluated in four groups according to Kern et al recommendations (Crit Rev Oncol/Hematol 2005). Results: The diagnostic samples in AML patients for FCM were obtained from bone marrow (n=133) or peripheral blood (n=33). The median value of immature cell ratio was 63 % (21%-98%). We detected leukemia associated IP(LAIP) in one and/or more than one categories at the diagnosis (Table): In summary: The 55 of the diagnostic samples, were positive for cross-lineage expression of lymphoid antigen, 148 show asynchronous expression of antigens, 155 present a lack of myeloid antigen expression and 111 have had myeloid antigen overexpression. The most frequent LAIP in the diagnosis was the persistency of immature myeloid antigen during the maturation (33. 4%) or the lack of expression of myeloid antigen (34. 5%). Only 73 patients were able to be analyzed for MRD on day of 14th to 54th following remission induction(RI) therapy. We could not evaluate 17 patients (23%) for MRD at the follow-up in which hematological remission was not achieved after a RI therapy. Eighty-nine percent of 56 patients in the remission had any LAIP permitting for the evaluation of MRD. The amount of MRD was 0.1% to 10% in different follow-up(F/U) periods. We observed relapse within 6 months in 36% of 55 patients having MRD. Conclusion: We found less myeloid-antigen associated abnormal IP(7. 65%) in the diagnostic samples.
Compared with the previously published studies (10-24%), however, the ratio of leukemic IP was detected as 92.6%. Evaluation of MRD with regular F/U of FCM events may allow us to predict the probability of relapse.

<table>
<thead>
<tr>
<th>Classification</th>
<th>LAIP</th>
<th>The detected parameter</th>
<th>%</th>
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</thead>
<tbody>
<tr>
<td>Cross-lineage</td>
<td></td>
<td>CD2+ , CD19+, CD4+, CD7+ etc</td>
<td>61</td>
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<tr>
<td>(n=55 patients)</td>
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<tr>
<td>Asynchronous</td>
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<td>CD117+/34+/33+±13+ -CD11b+/ CD117+/CD34+ -CD15+/ CD33+±CD19+ etc</td>
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<td>(n=148 patients)</td>
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<td>Lack of expression</td>
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<tr>
<td>Overexpression</td>
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<td>CD11b+/CD117+/CD34+/+ -HLA- DR+ +/CD33+/CD34+ etc</td>
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<td>(n=111 patients)</td>
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<tr>
<td>Total</td>
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</table>

**Early Recurrence of an AML-M4 Case Showing Normal Hematologic Parameters Presenting with Solitary Eye Involvement**

Mesut Ayer, 1Makbule Ulusoy, 1F. Aylin Ayer, 2Yesim Gurkan, 2Hikmet Feyizoglu, 2Namik Yigit, 2Zekai Kuyubasi, 1Arzu Karacevik

1Haseki Training and Research Hospital, 4th Internal Medicine Clinic, Istanbul, Turkey 2Haseki Training and Research Hospital, 2nd Internal Medicine Clinic, Istanbul, Turkey

Peripheral blood and bone marrow assessments of a 50 yrs old male patient presenting with leucocytosis, anaemia and thrombocytopenia in May 2006, revealed 90% of blasts some of which included Auer bodies. Following flowcytometric and cyto genetic(t[15;17],15) evaluation the patient was diagnosed as “acute myelomonocytic leucemia” (AML-M4). The patient received 3+7 treatment for remission induction. After attaining remission, 3+7 treatment was repeated in order to achieve consolidation. The patient did not have a suitable HLA donor so discharged from the hospital with a consolidation treatment schedule including high dose of Cytarabine (HiDAC). At the end of the second month ptosis developed at the left eye. Blood count analysis was within normal range. While the patient was waiting for hospitalization ptosis developed at the other eye together with external divergence of the left eye. At the time of hospitalization blood count analysis was repeated and found normal. Bone marrow aspiration analysis revealed 20% of blasts. Cell count, smear and flowcytometric evaluation of cerebrovascular fluid was normal. His eye fundus examination was normal yet involvement of 4th cranial nerve (n. trochlearis) was considered. Orbital MR imaging was normal. Development of an early recurrence was suspected so FLAG-IDA treatment regimen and radiation therapy on the orbital zone was applied. His control bone marrow investigation at the 28th day revealed remission and eye findings totally regressed. FLAG-IDA treatment was repeated for the second time in order to get consolidation. The patient was discharged from the hospital and hematology outpatient clinic appointments were organized. Low Dose of Cytarabine (LD ARA-C, 20mg/m2/day, 8 days a month) treatment was scheduled but the patient presented with leucocytosis and peripheral blood analysis revealed recurrence at the third week visit. This case report points out the possibility of early recurrence with eye involvement while hematologic parameters are normal.

**Molecular Hematology**

Ref. No: 40 Abstract No: 17

**Differential Expression of 16 Apoptosis Related Genes in Vitamin D-Induced Differentiation of HL-60 Cells.**

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1Kocaeli University Faculty of Medicine Department of Medical Biology, Kocaeli, Turkey 2Kocaeli University Faculty of Medicine Department of Medical Genetics, Kocaeli, Turkey

Using a quantitative real-time PCR (LightCycler), we analyzed 16 genes (Bcl-2, Bcl-xl, Mcl-1, Bik, Caspase 6, Caspase 7, Cytochrome-c, TNFR1, Myc, TGF-beta, JNK1, p38MAPK, p21, p27, Cdk2, Cyclin E) for changes in expression associated with the apoptosis of human promyelocytic leukemia HL-60 cells induced by 1alpha,25-dihydroxyvitamin D3 at various time points 18, 48 and 72h. Cells were cultured and RNA samples were isolated. Relative quantification data obtained from PCR products of cDNA portions. We did not find distinct down or up-regulated expression profiles at these time points. Findings suggest that there are not clear apoptotic signals in early phases of differentiation and the genes involved in vitamin D-induced apoptosis of HL-60 cells would be more clearly visible after the terminal differentiation process. Apoptosis and cell cycle analysis could be performed in this experimental setting, using quantitative real-time PCR.

Ref. No: 94 Abstract No: 18

**Circulating Endothelial Cells in Leukemic Patients Who Have CMV Antigenemia.**

Ilknur Kozanoglu, 1Can Boga, 2Hakan Ozdogu, 3Oktay Sozer, 1Ercan Maytalman

1Baskent University Medical Faculty Physiology Department, Ankara, Turkey 2Baskent University Medical Faculty Hematology Department, Ankara, Turkey 3Baskent University Adana Hospital Hematology Research Laboratory, Adana, Turkey

Human cytomegalovirus (HCMV) pathogenesis is dependent on the hematogenous spread of the virus to host tissue. While data suggest that infected monocytes are required for viral dissemination from the blood to the eye involvement with presentation of ptosis and strabismus.
host organs, infected endothelial cells are also thought to contribute to this key step in viral pathogenesis. Infection of endothelial cells promotes the increased surface expression of cell adhesion molecules (intercellular cell adhesion molecule 1, vascular cell adhesion molecule 1, E-selectin, and platelet endothelial cell adhesion molecule 1), which were necessary for the recruitment of naive monocytes to the apical surface of the endothelium and for the migration of these monocytes through the endothelial cell layer. We aimed to quantitate endothelial progenitor (EPCs) and circulating endothelial cells (CECs) by using flow cytometry in CMV pp65 positive leukemic patients. Detection of CMV pp65 matrix protein in peripheral blood leukocytes by using immunofluorescence assay (Chemicon Internation, a Serologicals Company). All slides are preperared 200,000 cells and investigated on fluorescence microscope (Eurostar, Euroimmun GmbH, Lübeck, Germany). At this time a panel of monoclonal antibodies, anti CD146 FITC, anti CD 144 PE, anti CD34 ECD, anti CD117 PC5 were used to enumerate CECs and EPCs in CMV pp65 positive patients. Flow cytometric measurement were performed with a FACS calibur flow cytometer (Coulter Epics XL- MLC, Beckman Coulter, Florida, USA) equipped with a 15 mW air-cooled 488-nm argon ion laser. Data were analyzed by using EXPO 32 ADC software. We analyzed CECs and EPCs by flow cytometry in CMV pp65 positive 3 patients (one AML M0, one chronic leukocytic leukemia and one hairy cell leukemia. CMV pp65 positive (40 cells) in AML patient, CECs 30. 540/mL and EPCs 14000/mL. One week later, after antiviral therapy, 100 CMV pp65 positive cells detected while CECs number was 7860/mL and EPCs 4350/mL. On the subsequent week, 5 and 2 CMV positive cells detected on the fluorescence microscope and CECs and EPCs number was decreased. Similar results was shown in CMV positive chronic leukocytic leukemia and one hairy cell leukemia patients (Table1). During the past decade an increasing population of immunosuppressed individuals has resulted in a resurgence of CMV as a major pathogen. Induced immunosuppression has occurred more frequently via chemotherapy and transplant regimens. Detection of blood leukocytes is closely associated with the clinical manifestations of CMV disease and is useful in the diagnosis of CMV infection. We concluded that; CECs and EPCs identification method in peripheral blood of patients with pathophysiological manifestations involving endothelial damage that are different from those caused by CMV infections, can be performed. Our observation supported the idea that CECs and EPCs quantit

<table>
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<th>CMV pp65(+)</th>
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<th>CEC / ml</th>
<th>EPC / ml</th>
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ACUTE MYELOBLASTIC LEUKEMIA

Ref: No: 96 Abstract No: 19

COLONY FORMING ASSAY IN PATIENTS WITH AML M7

1Ilkennur Kozanoglu, 2Can Boga, 3Hakan Ozdengo, 3Ercan Maytalman, 3Nihan Aldirmaz
1Baskent University Medical Faculty Physiology
Department, Ankara, Turkey 2Baskent University Medical Faculty Hematology Department, Ankara, Turkey 3Baskent University Adana Hospital Hematology Research Laboratory, Adana, Turkey

The discovery of the colony-forming capacity of hematopoietic precursor cells 40 years ago has revolutionized experimental, diagnostic and therapeutic hematology. In addition to quantitative abnormalities, cultures can reveal disease-specific diagnostic growth patterns. They are help in the staging of some hematological disease. In untreated AML, cultures of bone marrow show either no growth, or growth of leukemic cells either in clusters or as single cells. Leukemic growth can be discrete or abundant; single cells in leukemic clusters sometimes resemble either neutrophils, eosinophils, macrophages or immature erythroblasts according to their line of origin. The typical in vitro growth pattern in AML was observed soon after the discovery of the hematopoietic stem cell. Its association with an unfavorable prognosis, its disappearance in remission and its value in the prediction of relapse are well documented. On the other hand, absence of leukemic growth is an acknowledged good prognostic sign. The aim of our study were to evaluate endogenous colony formation in AML M7 patients. The bone marrow mononuclear cells of 2 patients with AML M7 were cultured by cytokine free methyl cellulose media (Methocult TM GF H4230 – StemCell Technologies, Canada) for endogenous colony evaluation. removed 100 μl blood, counted initial nucleated cell by adding 3% acetic acid and using hemacytometer or counted directly in automated cell counter. Firstly, it was added 1/3 of ficoll and then added 2/3 of blood on the top of ficoll gently. It was santrifugated 400xg for 30 min. It was removed layer of mononuclear cells between sera and ficoll. It was added IMDM containing 2% FCS on the mononuclear cells and resuspended them and santrifugated 400xg for 10 min. It was decanted the supernatant repeat the step above. Decant the supernatant and then resuspended mononuclear cells in the way that cell count is 1-2,5 x 106 by adding IMDM containing 2% FCS (in order to get cell count, perform the step mentioned on the top). It was thawed an alliquot overnight between 2-8 °C. Added 0,3 ml of resuspended cell prepared before. Last concentrati

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progenitors reappeared. AML M7 can be diagnosed easily by flow cytometry.

Ref. No: 99 Abstract No: 20
THERAPEUTIC LEUKOCYTAPHERESIS: THE APPLICATION OF RESULTS OF SINGLE CENTER
1 İlkınur Kozanoglu, 2 Can Boga, 3 Hakan Ozdogu, 4 Mahmut Kural
1 Baskent University Medical Faculty Physiology Department, Ankara, Turkey 2 Baskent University Medical Faculty Hematology Department, Ankara, Turkey 3 Baskent University Adana Hospital Apheresis Unit, Adana, Turkey

Therapeutic leukopheresis is one of the recommended treatment modalities in hyperleukocytosis - leukostasis, bleeding disorders related to some subtypes of leukemia and tumor lysis syndrome after chemotherapy. Leukopheresis should be performed if leukocyte count is over 100x10^9/L in AML and ALL; and over 300x10^9/L in CML associated with clinical findings of leukostasis. Therapeutic apheresis procedures were performed by using COBE Spectra (Cobe, Lakewood, CO, USA) working continuous blood flow. Between December 2003 and April 2007 total 76 leukopheresis procedures were performed in 43 patients (16 female, 27 male). There were followed up 23 patients with acute leukemia and 20 patients with chronic leukemia. Forty to fifty percent of reduction in leukocyte count in all procedures. All of the patients completed the procedure without any severe adverse events related to the procedure. We did not any severe complication. In certain mostly emergent conditions bedside therapeutic leukopheresis is helpful. In conclusion, however a instutionan a dedicated staff and a mobile cell separator for this application are needed.

Ref. No: 111 Abstract No: 21
SUPPORTIVE CARE BY GRANULOCYTE COLONY-STIMULATING FACTOR IN HYPOPLASTIC ACUTE MYELOGENOUS LEUKEMIA
Can Boga, Hakan Ozdogu, Mahmut Yeral, Ebru Kızılkılık, Mutlu Kasar
Baskent University Faculty of Medicine, Department of Hematology, Adana, Turkey

Oligoblastic leukemia mostly occurs in elderly patients and appears to proliferate slowly and be characterized by a maturation failure. The standard use of chemotherapy generally is ineffective. Moreover in the majority of the elderly patients, chemotherapy has substantial toxicity. Some reports have shown that granulocyte-colony-stimulating factor (G-CSF) has anti-leukemic activity in oligoblastic leukemia. Herein, two elderly patients with oligoblastic acute myeloid leukemia received subcutaneously G-CSF in addition to supportive care. The patients had no multilineage dysplasia and high-risk karyotype. During G-CSF treatment (filgrastim 300 μg/day) absolute neutrophil counts (>0.5x10^9/L, for two patients) rose to 1x10^9/L, and absolute platelet counts (<30x10^9/L, for two patients) increased of 20x10^9/L or more). A marrow blast clearance (<5%) was not observed in these patients. They were transfusion independence for the first month. However, on the second month of initiating therapy, leukemic cells increased in the peripheral blood. One patient died during treatment after a 6 months due to sepsis. One patient was still on treatment with combination standard induction chemotherapy. In acute myeloid leukemia, G-CSF has been used to reduce the duration of neutropenia after induction or consolidation therapy. Recently, spontaneous complete remission of acute myeloid leukemia has been reported after treatment with G-CSF alone without chemotherapy. The mechanism of action of filgrastim probably is the selective stimulation of normal residual marrow. Hence, filgrastim can be accepted as a part of the supportive care. Our observation supported this idea.

Ref. No: 114 Abstract No: 22
EMA AS SALVAGE REGIMEN IN PATIENTS WITH REFRACTORY/RELAPSE AML: A SINGLE CENTER EXPERIENCE
Pervin Topcuoglu, Sinem Civriz, Bilge Ceydilek, Meltem Kurt Yüksel, Sule Mine Bakanay, Onder Arslan, Taner Demirer, Arkın Uysal, Meral Bekşac, Günhan German, Mutlu Arat, Muhit Ozcan
Ankara University, School of Medicine, Ankara, Turkey

In this retrospective study, we evaluated that the effect of EMA as salvage regimen on the short- or long-term outcomes in 44 patients with refractory or relapse AML between March 2003-June 2006. Patients: Median age was 32 years (17-57 years). The ratio of male/female was 27/17. The distribution of the patients according to FAB classification as follows: AML-M0: 1; M2: 7; M4-M5: 20 and M6: 2; biphenotypic leukemia: 2 and secondary AML: 3. EMA as salvage regimen for induction treatment was given to 16 patients who underwent primary refractory of the induction regimen and 22 patients of first relapse (early relapse within 12 months: 22; Late relapse: 6). EMA treatment protocol: Mitoxantrone, iv daily 12mg/m2/day on day 1 to day 3; Etoposide, iv daily 200mg/m2 on day 8 to day 10; medium dose cytosine arabinoside, continue infusion 500 mg/m2/day from day 1 to day 3 and from day 8 to day 10. G-CSF (Neupogen) daily 5μg/kg by sc route had been given from 12th day until absolute neutrophil count reached above 0.5x10^9/L in consequent 3 days. Results: Total six patients died of bacterial or fungal infection at the aplasia period during the induction of EMA. When we evaluated the response in the rest of the patients (n=38), complete remission with EMA was observed in 57.8% of the patients (n=22) (figure 1). After the EMA treatment, 13 patients received only chemotherapy (n=7) and/or underwent autologous hematopoietic cell transplantation (allo-HCT) as a consolidation treatment (n=6). Seventeen of 22 patients being in CR after EMA have currently been in disease-free survival (Mean 22.8 months; range: 15-730.6). In 13 of 16 patients with unresponsive of the EMA protocol, EMA±CsA (n=12) or other salvage regimens (n=2) were used again for reinduction, and CR was obtained in only 6 patients (46%). These 6 patients are still alive in present time, but one of them has relapse disease. When we analyzed the patients as a whole median follow-up period from the diagnosis was 13.2 months (95% CI: 12.09-14.3 months) and the probability of two-year overall survival (OS) was 47.4±9.1. The probability of two-year DFS and OS after EMA treatment were 23.4±8.4% and 40.4±10.1%, respectively. When we evaluated the patients in two groups, primary refractory or relapse as indication of EMA use, OS from the diagnosis was significantly shorter in primary refractory group than the relapse group (median 16.8 months vs 58.3 months, p=0.006). After the EMA treatment, these two groups did not lead to a statistical difference on OS (26.7±13.8 months vs 10.7±7.6 months, p=0.326). In conclusion, we observed that the use of EMA as salvage regimen had a
positive effect on DFS and OS from the diagnosis in both primary refractory and relapse patients.

The disease course and survival as the indication of EMA use

Abbreviations: CR: Complete remission; Allo-HCT: Allogeneic hematopoietic cell transplantation

Ref: No: 116 Abstract No: 23

ACUTE MYELOID LEUKEMIA PRESENTING AS ACUTE INFERIOR MYOCARDIAL INFARCTION
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A 27-year-old man was admitted with retrosternal pain, and sweating lasting 2 hours. Electrocardiography showed signs of acute inferior myocardial infarction. The patient did not report any symptoms or any complaint before. He had no history of cardiac disease and he was not smoking. His arterial pressure was 110/70 mmHg and cardiac pulse was 98 per minute. The physical examination was normal. Laboratory tests showed a white blood cell count of 8.4x10^9/L (84% monocyte), platelet count of 106x10^9/L and hemoglobin of 11 g/dl. Blood chemistry showed elevated lactic dehydrogenase (280 IU/L), creatine phosphokinase (1120 IU/L), glutamic-oxaloacetic transaminase (52 IU/L) and troponin I (9 IU/L), but no elevation in aPTT, PT and D-dimer. triglyceride, total cholesterol, protein C, protein S and antithrombin 3 was normal. Coronary angiography was performed. This showed thrombotic occlusion of the right coronary artery. Aspiration of thrombus and local t-PA was applied in coronary artery angiography sessions. It was observed normal perfusion of the artery after the procedure. Because of the cytopenia and monocytosis; peripheral blood smear and bone marrow examination was performed. The patient’s peripheral blood smear, bone marrow films and flow cytometric examination revealed acute myeloid leukemia FAB M0. Various conditions, such as leukemic infiltration into the myocardium, occlusion of a major coronary artery by leukemic thrombus, effects of the antileukemic therapy (such as chemo therapy, or radiation therapy), hypercoagulable state or hemorrhages in the myocardium or intima of a coronary artery can cause a myocardial infarction in leukemia. In conclusion the reason of coronary artery occlusion in our patient of acute myeloid leukemia remains speculative.

Table 1. Patient Characteristics

<table>
<thead>
<tr>
<th>Age (year, median/range)</th>
<th>53/19-77</th>
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<tbody>
<tr>
<td>Sex (F/M)</td>
<td>15/20</td>
</tr>
<tr>
<td>WBC(μl,median/range)</td>
<td>28 100/310-457 000</td>
</tr>
<tr>
<td>Hb( g/dl, median/range)</td>
<td>8.1/4.0-13.0</td>
</tr>
<tr>
<td>PLT(μl,median/range)</td>
<td>35 000/7 000-520 000</td>
</tr>
<tr>
<td>LDH(U/l,median/range)</td>
<td>352/195-2477</td>
</tr>
<tr>
<td>FAB subtype</td>
<td></td>
</tr>
<tr>
<td>M0-M1</td>
<td>14 (%40)</td>
</tr>
<tr>
<td>M2</td>
<td>12 (%34.3)</td>
</tr>
<tr>
<td>M3</td>
<td>4 (%11.4)</td>
</tr>
<tr>
<td>M4</td>
<td>3 (%8.8)</td>
</tr>
<tr>
<td>M5</td>
<td>2 (5.7)</td>
</tr>
<tr>
<td>CD34(+)</td>
<td>28 (%74.28)</td>
</tr>
<tr>
<td>Organomegaly (lymphadenopathy, hepatomegaly, splenomegaly)</td>
<td>13 (%37.14)</td>
</tr>
<tr>
<td>CR</td>
<td>11(%57.89)</td>
</tr>
</tbody>
</table>

Table 2.

<table>
<thead>
<tr>
<th>UPAR</th>
<th>p&lt;0.05</th>
</tr>
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<tbody>
<tr>
<td>Age</td>
<td>p&lt;0.05</td>
</tr>
<tr>
<td>Sex</td>
<td>p&lt;0.05</td>
</tr>
<tr>
<td>WBC</td>
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<tr>
<td>Hb</td>
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<td>PLT</td>
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<td>Organomegaly</td>
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<td>CD34</td>
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<td>CR</td>
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</table>

Ref: No: 120 Abstract No: 24

EXPRESSION OF UROKINASE PLASMINOGEN ACTIVATOR RECEPTOR IN ACUTE MYELOID LEUKEMIA
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Background: Urokinase plasminogen activator receptor (UPAR) is a membrane protein responsible for plasmin expression on cells facilitating cellular extravasations and tissue invasions. Aims: To determine UPAR expression incidence rate by flowcytometry method from patients with acute myeloid leukemia (AML). Methods: This study included 35 patients with AML. A variety of clinical and biological parameters, including phenotype have been examined for potential value in predicting treatment response. The patients characteristics are presented in Table 1. We evaluated the correlation between UPAR and age, sex, organomegaly, CD34, complete remission (CR) rate. Results: The correlation values are presented in Table 2. We have found significant correlation between organomegaly and UPAR. The highest UPAR expression was seen in cases with AML M4 and M5. Summary / Conclusions: We observed a high expression of the UPAR in myelomonocytoid subtypes (M4, M5) in AML. Monocytoid differentiated leukemias are characterized by high rates of extramedullar disseminations. UPAR binding to the UPAR initiates the conversion of plasminogen to the protease plasmin, which mediates the extravasation of cells through the endothelium by proteolytic cleavage of endothelium associated adhesion molecules. UPAr expression could play a role in adhesion, migration, and metastasis of leukemic cells. Therefore UPAR is correalted with organomegaly in AML cases.
NEPHROTIC SYNDROME AS THE FIRST MANIFESTATION OF ACUTE MYELOGENOUS LEUKAEMIA: CASE REPORT

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Introduction: The hematological malignancies associated with nephrotic syndrome are mainly hodgkin’s and non-hodgkin’s lymphomas and chronic lymphocytic leukemia. Acute myelogenous leukemia (AML) has rarely been described in association with nephritic syndrome. We report a rare case of Acute Myelogenous Leukemia who presented with nephrotic syndrome. Case report: A previously healthy 62-year-old man was admitted in nephrology ward because of generalized developing pitting edema during last month. Simultaneously, he had generalized itching and urticaria, polyuria, polydypsia and low grade fever but had no history of weight loss, anorexia and sweating. In laboratory tests he had proteinuria above 3.5 g/day. Because of anemia, hematology consultation was done. In peripheral blood, there were myeloblast cells in the circulation at a ratio of 15%. Bone marrow aspiration confirmed a diagnosis of AML, showing hypercellular bone marrow with 80-90% leukemic cells, increased M/E ratio, myeloblast (immature cell, fine chromatin, cytoplasmic blob) and these abnormal elements: >50% and mature cell about 10%. Unfortunately, we hadn’t renal biopsy as a consequent of patient illness and thromboastenia. He received induction chemotherapy, which led to a complete remission and decreasing urinary protein excretion during chemotherapy and no proteinuria at the end of it. Now the patient has received second course of consolidation therapy and remained in complete remission, with no physical and laboratory evidence of proteinuria. Conclusion: It can be concluded that nephrotic syndrome may be additionally associated with AML. In some cases, there is a direct causal effect of the leukemic process on renal function or even pathology, while in others it is exerted indirectly via other complications of the malignancy or the treatment. KEY WORDS: AML, nephrotic syndrome, case report

CHRONIC MYELOID LEUKAEMIA

ORAL AND CUTANEOUS LICHENOID REACTION SECONDARY TO STANDARD DOSE IMATINIB: A CASE REPORT

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Chronic Myeloid Leukemia (CML) is a clonal myeloproliferati disorder which is the first human malignancy to be associated with a specific genetic lesion, the Philadelphia chromosome, carrying BCR-ABL oncogene. Imatinib (Glivec) is the first molecularly targeted drug developed for CML and has achieved a remarkable success. Cutaneous side effect with this treatment are common (especially with High Doses) but lichenoid reaction on drug eruption is rare. Only a few cases of oral and/or cutaneous lichenoid reaction secondary to imatinib have been reported. We present a 42-years old woman with approximately one year history of CML. She was treated with standard dose Imatinib (400mg). Three months after the beginning of this treatment, while using the drug, grey-violaceous plaques with a reticular pattern resembling oral lichen planus on both cheek mucosal surfaces and a disseminated cutaneous eruption appeared on the trunk, legs and arms and composed of dark purple, pruriganeous papules suggestive of lichen planus. The cutaneous and the oral mucosal biopsy confirmed the diagnosis of lichen planus. It is believed that this patient developed imatinib-induced lichenoid eruption; this relationship with therapy rather than the underlying diseases rules out a paraneoplastic reaction and idiopathic lichen planus. Key words: CML, Imatinib, lichenoid eruption
Chronic Lymphocytic Leukemia (CLL) is the most common form of adult leukemias. Many factors have been identified to determine prognosis in CLL over the past 10 years. Importance of genetic studies has advanced in last few years. Whereas conventional techniques reveals genetical defects in only 40-50% of cases, new molecular techniques such as fluoresan in situ hibridisation (FISH) has provided very important improvements in detecting chromosomal defects. Cytogenetic abnormalities seen in CLL are deletion 13q14 (55% of all cases), deletion 11q (18 %), trisomy 12q (16 %), deletion 17p (7 %) and deletion 6q (6 %). Deletion 11q is related to lymphadenopathy and rapid disease progression. Deletion 17p predicts treatment failure with alchylating agents such as fludara-bine and short survival. In CLL cases without Immune globulin variable heavy chain mutation short survival times and rapid disease progression are predicted. Whereas survival times have been found the shortest, only 32 months, in patients with deletion 17q, the longest, 133 months, in cases with deletion 13q. Expression of CD38 on 6q (6 %). Deletion 11q deletion in one, normal karyotype was found in 11 %, hypodiploidy in 22 of patients and among these patients, bone marrow involvement pattern, serum lactate dehydrogenase (LDH) and beta2 microglobulin (β2M) levels). We obtained the file records of our CLL cases and collected cytogenetic analysis results of 33 CLL patients who had bone marrow aspiration and biopsy between 2002 and 2006. Clinical stages of patients were Rai stage 0 (13 patients, 39,3 %), stage I (4, 12,1 %), stage II (7, 21,2 %), stage III (one, 3 %) and stage IV (7, 21,2 %). One patient (3 %) had transformation to Richter Syndrome from stage II. There were at least one constitututional symptom in 2 patients (15,3 %) with stage 0, in 2 patients (50 %) with stage I, in 3 patients (28,5 %) with stage II, in none of patients with stage III and in 5 patients (71,4 %) with stage IV. Lymphocyte doubling times of 7 (21,2 %) patients were below 12 months. Serum LDH levels were more than normal in 13 % (39) patients. Serum β2M levels were more than normal in 20 (60 %) patients. Bone marrow biopsy was performed in 22 of patients and among these patients, bone marrow involvement patterns were nodular in 12 patients (54,5 %), diffuse in 8 patients (36,3 %), nodulary and diffuse in 2 patients (9,2 %). Conventional cytogenetic analysis was used for all the patients. Among the 18 patients only detected with conventional technique, there were normal karyotype in 8, no reproduction in 5, hypodiploidy in 4, and 7q deletion in one patient. Fifteen patients were evaluated with FISH and 13q deletion was detected in 3, 11q deletion in one, normal karyotype was found in 11 of CLL patients.
ting factor and erythropoietin were administered. After 2 months, her clinical findings did not improve, only neutrophils increased and splenic irradiation applied. Splenomegaly regressed (5 cm palpable). CSA and growth factors treatment continued. After six months, splenomegaly (1 cm palpable) and hepatomegaly (2 cm palpable) and CBC values improved. Leucocytes: 13400 cells/mm3, hemoglobin: 10.6 g/L, platelets: 198000 cells/mm3. Differentiation gave 88 % of neutrophils, 10 % of lymphocytes and 2 % of monocytes. CSA dose were tapered and discontinued. Because of her clinical picture reappeared, CSA treatment restarted but renal toxicity developed and CSA treatment stopped. She became transfusion dependent. She has administered first cycle of treatment on oral fludarabine 40 mg/m2 for 5 days. After four weeks of treatment, organomegaly regressed and CBC normalized. Leucocytes: 6260 cells/mm3, hemoglobin: 14.1 g/dL, platelets: 122000 cells/mm3. Differentiation gave 82 % of neutrophils, 14 % of lymphocytes and 4 % of monocytes. Second cycle of fludarabine treatment administered. But she developed headache and multipal cranial neuropathy including n. opticus, n. oculomotorious, n. trigeminalis and n. vestibulocochlearis, and her cranial MRI was normal. This multipal cranial neuropathy may depend on oral fludarabine treatment. It should keep in mind that fludarabine may cause severe multipal cranial neuropathy.

SEVERE NEUROTOXICITY OF CLADRIBINE IN A PATIENT WITH HAIRY CELL LEUKEMIA: A CASE REPORT

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Hairy cell leukemia (HCL) is an indolent chronic B cell lymphoproliferative disorder involving the bone marrow and spleen. Patients with HCL present with peripheral blood cytopenias, organomegaly and circulating hairy cells. Past a few years cladribine purine analogue has an important role with %90-100 remission rates in HCL treatment. We presented a patient with hairy cell leukemia who developed severe neurotoxicity after cladribine infusion. A 52 years old male patient admitted to the hospital with fatigue, weakness and weight loss continuing for 2 months. On his physical examination; blood pressure 100/60 mmHg, fever: 36 C, heart rate: 72, cardiovascular and respiratuary systems were normal. Spleen was palpable 10 cm below the costal margin and hepatomegaly determined. Laboratory findings were as follows: complete blood count; hemoglobin: 11g/dL; hemocrit: 32 %, wbc: 2500 /mm3, plt: 56000/mm3, erythrocyte sedimentation rate: 10 mm/h. In CT scans: 2 cm diameter bilateral axillar, 4 cm diameter cheekal, peripancreatic, paraaortic and paracaval lymphadenomegalias were observed. We also determined spleen enlargement (long axis was 22 cm) and hepatomegaly before treatment. The diagnosis was established by examining bone marrow biopsy with immunohistochemical stain. Bone marrow biopsy was examined in Ege University Pathology Department. Hairy cell leukemia infiltration was found, CD20 (+), TRAP (+), CD3 (-), CD68 (-) and reticulin grade was 2 respectively. Our case received cladribine 0.1mg/kg/day, 7 day duration infusion. Cladribine therapy generally has minimal acute or subacute adverse events. Besides this, moderate bone marrow inhibition is the most common adverse event. One week after the treatment the patient developed neutropenic fever, pulmoner infection, weakness in both legs, short term memory losts, and mislaid ability to walk. Then he received appropriate antibiotic and antifungal therapies. Cranial MRI and EEG was normal but EMG showed early stage axonal degeneration and segmental demyelination which is significant in lower extremities. These neuropathic findings were interpreted a neurotoxicity of cladribine by the neurologists. 6 weeks after the treatment his general condition became well and laboratory findings regressed. Thorax and abdominal CT scans were all normal. Almost he regained his ability to walk and muscle strength. Our case is in remission hematologically but still going on neurology controls. When literatures have reviewed severe neurotoxicity was seen rarely. In National Cancer Institute group 932 cases were treated with cladribine and 26% of these had developed moderate neurotoxicity. Grade 3-4 neurotoxicity had developed only in 10 patients in this series. We approved to present this case which developed opportunistic pulmoner infection, hemolytic anemia and serious neurotoxicity after standard dose treatment.

POSSIBLE PNEUMOCYSTIS JIROVECI INFECTION IN AN CYTOMEGALOVIRUS POSITIVE CLL PATIENT

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Chronic lymphocytic leukemia (CLL) is the most common form of leukemia in Western countries and mainly affects elderly individuals. Infections are major cause of morbidity and mortality in patients with chronic lymphocytic leukemia. Predisposition to infection in CLL is mediated through various abnormalities including both the impairment of humoral and cellular immunity and further immunosuppression related to the therapy of CLL. Streptococcus pneumonia, staphylococcus aureus, streptococcus pyogenes and herpes zoster-varicella virus are most common infection agents. On the other hand, patients treated with purine analogues apparently have an increased incidence of infection with other opportunistic organisms such as cytomegalovirus, listeria monocytes and pneumocystis jiroveci. We are presenting a patient with CLL associated with CMV and pneumocystis jiroveci infection. A 63 year old man with 3 year history of CLL was treated with fludarabine and cyclophosphamide. Because of development of autoimmune hemolytic anemia 2 weeks before admission to our hospital, steroid therapy of 70 mg prednisolone was started. Ten days after second course of fludarabine and cyclophosphamide therapy, he presented with fever. The white count cell on admission was 2.8 x 109/L (%30 lymphocytes). Physical examination and chest radiography was normal. Empirical meropenem therapy was started. The patient was unresponsive. Test for CMV pp65 antigenemia was found to be positive. Ganciclovir was started. Five days after ganciclovir therapy, progressive dyspnea occurred. Computed tomographic examination of the thorax revealed consolidation and bilateral pleural effusion. The patient was not tolerated bronchoscopy and empirically high dose co-trimoxazole was added to therapy, the clinical condition of the patient was improved. Atypical double opportunistic infections may see in CLL patients who are treated with purine analogs. Association of these two infections should keep in mind in these patients.
PROMINENT PLEURAL AND PERICARDIAL EFFUSION DUE TO IMATINIB MESYLATE AFTER FIVE YEARS OF THERAPY

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Baskent University, Faculty of Medicine, Department of Internal Medicine, Hematology Division, Ankara, Turkey

The signal transduction inhibitor imatinib mesylate, is the superior first line therapy in Philadelphia chromosome positive chronic myeloid leukemia (CML) patients. Data obtained so far has shown that imatinib mesylate can induce hematologic and cytogenic remissions in CML patients in all stages of disease. The drug is generally very well tolerated. The most common side effects are mild nausea, myalgias, edema and diarrhea. The manufacturer suggests, especially patients over 60 and with higher doses of imatinib mesylate unexpected weight gain should be carefully monitored. Severe fluid retention was reported 1%-2%. The mechanism of this side effect is not clear. In the literature there are few cases reporting pleural-pericardial effusions after long term therapy. We describe this uncommon complication in a patient without progression of disease and at the dose of 400 mg imatinib mesylate. A 73 year of female patient admitted to our hospital with chest pain and dyspnea. She was diagnosed as Ph-positi-
ve CML 5 years ago and she was using imatinib mesylate since then. Before this admission there had been no signs of fluid gain ever. This time she had moderate pericardial effusion (no tamponade) and large pleural effusion. There are no symptoms of infection and no signs of infiltration and pulmonary embolism on the thoracic computerised tomography. Total amount of 1950 cc pleural fluid was drained on two occasions. Both were transudative, ADA (adenosin deaminase) level was normal, microscopy showed lymphomononuclear cells and culture was negative. Cytologic evaluation of the liquid did not show any myeloid or blastic cells. Ejection fraction and wall motions of the hearth was normal. No immature myeloid cells were seen on peripheral blood smear and there were no blastic cells in bone marrow sample. Imatinib treatment was discontinued and furosemide was started. Five days after pericardial effusion dissolved. Resolution of pleural fluid lasted four weeks. After one month of clinic stability imatinib was restarted. Until then there is no sign of fluid retention and the patient is in hematologic remission. In the present case we excluded progression of disease and extramedullary leukemia infiltration. There were no signs of infection and heart failure and other commonly known etiologies for effusions were excluded too. After cessation of imatinib mesylate pericardial effusion dissolved in five days but pleural effusion remained a problem for four weeks. There are cases reports which steroids were used in the literature but for this case furosemide was the only agent used for therapy. To our knowledge this the only case with 400 mg dose after 5 years of therapy. This case reminds that there can always be a risk for serious effusion in these patients even with standart dose. Imatinib can be started again after effusions resolved and may not cause the same clinic ever again. These patients must always be followed for recurrence tough.

THE CLINICAL PRESENTATION AND DIFFERENT BCR-ABL TRANSCRIPTS IN CML

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1Akseniz University, School of Medicine, Antalya, Turkey
2Stileyan Demirel University, School of Medicine, Isparta, Turkey

The importance of different BCR/ABL transcripts in CML has not been explained exactly yet. It was reported that in almost all CML the breakpoints in the BCR gene are found within the M-bcr. It is also constituted two different transcripts named b3a2 and b2a2 both which encodes p210 protein. The breakpoints in the m-bcr region are rare in CML. Clinically p190+ CML patients have a clinical findings between CML and CMML. The aim of this study was to evaluate the association of BCR/ABL transcripts with the specific clinical features. Methods: We analysed the p210 and p190 BCR/ABL transcripts of 64 chronic phase CML patients and compared them according to their clinical (splenomegaly), laboratory findings (platelets (plt), hemoglobin (hb), white blood cell (WBC) counts) and Sokal score. Fifty one patients expressed p210 (79%), 10 patients had p190 (16%), 3 patients (5%) expressed both p210 and p190 transcripts. The three patients who had both p210 and p190 transcripts all had higher WBC counts and massive splenomegaly. We did the analysis after exclusion of these three patients. We couldn’t find significant difference between the study parameters and p210, p190 and both transcripts except WBC. Patients who expressed p210 transcript had significantly higher WBC count (p=0.028) than patients with p190. We then analysed according to b3a2 and b2a2 transcripts. There were 8 patients who expressed both transcripts. No significant difference was found between the p210 transcripts and the study parameters but all of the patients with both transcript had higher WBC count but not significantly. We had only 10 patients with p190 transcript (4 e1a2, 2 e1a3 and 4 both transcripts). WBC count was also tend to be higher in patients with e1a3 and platelets were slightly higher in e1a2 patients. No significant difference was found in Sokal score of patients. As conclusion patients who express p210 transcript tend to have higher WBC count in CML but, the different p210 transcripts do not have an effect on WBC, plt, hb and Sokal score in CML. The significance of p190 and both transcripts should be evaluated with more patients.

FLAG-IDA IN THE TREATMENT OF REFRACTORY/RELAPSED ACUTE MYELOID LEUKEMIA: SINGLE-CENTER EXPERIENCE

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Background: We evaluated the efficacy and toxicity profiles of the combination of fludarabine, high-dose cytosine arabinoside (AraC), idarubicin, and granulocyte colony-stimulating factor (G-CSF) in refractory/relapsed acute myeloblastic leukemia (AML) patients. Methods: Between February 2003 and April 2007, 24 AML patients were treated with FLAG-IDA (fludarabine 30 mg/m(2),
AraC 2 g/m(2) for 5 days, idarubicin 12 mg/m(2) for 3 days, and G-CSF 5 micro g/kg from day zero until neutrophil recovery). Results: Eighteen patients were in relapse after conventional chemotherapy including idarubicin 12 mg/m(2) for 3 days and cytarabine 200 mg/m(2) for 7 days, (3+7) protocols. Six patients had refractory disease (after 7 days of standard doses of cytarabine, 3 days of idarubicin) Recovery of neutrophils and platelets required a median of 18 and 21 days from the start of therapy. Complete remission (CR) was obtained in 16 of 24 patients (66. 7%), 8 cases (33. 3%) had resistant to this regimen (RD) and 3 of 24 (12. 5%) died during reinduction therapy: 2 due to sepsis cerebral and 1 due to cerebral hemorrhage. Fever >38. 5 degrees C was observed in 20 of 24 patients (83. 3%), 15 had fever of unknown origin (FUO) and 9 documented infections; 16 of 24 (66. 7%) developed mucositis and 6 of 24 (25%) had grade 2 WHO transient liver toxicity. After achieving CR, 4 patients received allogeneic stem cell transplantation, 4 were judged unable to receive any further therapy, and 3 refused other therapy. Eight patients are at present in continuous CR after a median follow-up of 11 months (range: 2-20). Conclusions: FLAG-IDA is a good choice in cases with refractory/relapsing acute leukemia for salvage chemotherapy. High efficacy and a low-toxicity profile are preferable properties of this regimen, and this regimen has been found to be useful for cytoreduction, especially in candidates for allo-SCT.

HODGKIN’S LYMPHOMA

Objective: Comparison of the two treatment options in refractory and relapsing Hodgkin lymphoma, BEACOPP and EVA chemotherapy protocols, in terms of efficacy and frequency of febrile neutropenia. Methods: Patients refractory to treatment and patient not responding to treatment who have been treated for Hodgkin lymphoma between 1999-2006 were studied retrospectively. Statistical analysis was performed using the EpiInfo version 6 Statcalc software. Chi-Square test was used. Results: A total of 42 patients were studied in the study. All of these patients were found to have advanced stage Hodgkin lymphoma (Stage 3 and stage 4 according to the Ann-Arbor staging system). All of the patients were prescribed 6-8 cycles of ABVD chemotherapy as the first line treatment. Data analysis revealed that the number of patients receiving BEACOPP chemotherapy was 24 (14 males, 10 females), mean age was 33. 2 (26-42), mean duration between the end of ABVD chemotherapy and occurrence of relapse was 17. 4 (6-29) months, and 6 patients were refractory of ABVD chemotherapy. On the other hand, number of patients receiving EVA chemotherapy was 18 (11 males, 7 females), mean age 31. 8 (24-43), mean duration from the end of ABVD chemotherapy and the occurrence of relapse was 19. 2 (7-27) months, and 4 patients were refractory to ABVD chemotherapy. In both treatment protocols, patients were evaluated after the administration of 3 cycles of chemotherapy. Results showed that complete remission was observed in 13 patients (54. 1%) receiving BEACOPP, and partial remission was observed in 6 patients (25%) receiving BEACOPP. Five patients, however, did not respond to treatment (20. 8%). In the patient group receiving EVA chemotherapy protocol, however, complete remission was observed in 4 patients (22. 2%), and partial remission was observed in 4 patients (22. 2%). Ten patients (55. 5%) did not respond to treatment. In both groups patients not responding to treatment were referred to peripheral blood autologous bone marrow transplantation. The treatment protocols of patients with complete and partial remission were continued for 3 more cycles. Patients were re-evaluated after a total of 6 cycles. Analysis of the results of the group receiving BEACOPP showed that complete remission was obtained in 15 patients (62. 5%), partial remission was obtained in 4 patients (16. 6%). In the patient group receiving EVA chemotherapy group, however, complete remission was observed in 5 patients (27. 7%) and partial remission was observed in 3 patients (16. 6%). Febrile neutropenia was established in 11 patients (45. 8%) receiving BEACOPP therapy, and 3 patients (16. 6%) receiving EVA therapy (Table: 1). Table: 1- Patient characteristics and response of chemotherapy protocols. Discussion: EVA and BEACOPP chemotherapy protocols are treatment protocols used in refractory or relapsing Hodgkin lymphoma.
Comparison of the response rates to the administration of these two treatment protocols showed that BEACOP

<table>
<thead>
<tr>
<th>Patient Demographics</th>
<th>EVA/BEACOP</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number</td>
<td>18/24</td>
</tr>
<tr>
<td>Female Patients</td>
<td>7/10 (73%)</td>
</tr>
<tr>
<td>Male Patients</td>
<td>11/14 (50%)</td>
</tr>
<tr>
<td>Complete Remission</td>
<td>5/15 (33%)</td>
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<tr>
<td>Partial Remission</td>
<td>3/4 (66%)</td>
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<tr>
<td>Nodular Sclerosis Type</td>
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</tr>
<tr>
<td>Lymphocyte-rich Type</td>
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</tr>
<tr>
<td>Mixed Type</td>
<td>6/6 (100%)</td>
</tr>
<tr>
<td>Lymphocyte Deficient Type</td>
<td>5/8 (62%)</td>
</tr>
<tr>
<td>Stage 3</td>
<td>8/10 (80%)</td>
</tr>
<tr>
<td>Stage 4</td>
<td>10/14 (71%)</td>
</tr>
<tr>
<td>Number of patients with febrile neutropenia</td>
<td>3/11 (27%)</td>
</tr>
</tbody>
</table>

Ref: No 76

Abstract No: 38

HOEODGKIN’S DISEASE: A RETROSPECTIVE ANALYSIS OF 103 PATIENTS FROM A SINGLE REFERRAL CENTRE

1Osmangazi University Faculty of Medicine, Haematology Department, Eskisehir, Turkey 2Osmangazi University Faculty of Medicine, Biostatistic Department, Eskisehir, Turkey

A retrospective study was carried out in a group of 103 patients with Hodgkin’s disease (HD), 38 females and 65 males, ages 16-82 years (median 39 ± 15.54), who were treated and followed up in the period between 1986 and 2006 at Hematology Department of Osmangazi University Medical School. The clinical parameters used were sex, age at diagnosis, career, living place, type and number of nodal areas involved (peripheral, mediastinal or abdominal). Laboratory parameters considered included hemoglobin, absolute number of lymphocytes, ESR, serum levels of LDH, AST, ALT, GGT, ALP, fibrinogen and albumin levels, viral markers, blood type, immunoglobulin G, A, M and beta-2 microglobulin levels in addition to throracal-abdominal computed tomography findings, histologic type, stage and follow-up time. Of patients; 25.8% were housewives, 17.2% were workers, 16.1% were students, 16.1% were officers, 10.8% were retired, 7.5% were free-workers, and 6.5% were farmers. 83% of the patients were living in cities while 17% were from countries. Peripheral lymph nodes (cervical, axillary and/or inguinal) were the most commonly involved location. At advanced-stage (III/IV) ESR, LDH, ALP, GGT levels and hypogammaglobulinemia were increased while levels of hemoglobin and albumin were decreased significantly. Patients with advanced-stage were also older than patients with early-stage (I/II). In patients with advanced age sedimentation rate and beta-2 microglobulin were increased and albumin was decreased significantly. Patients in whom abdominal involvement was documented; there was a significant increase in sedimentation rate, beta-2 microglobulin and levels of LDH and ALP while hemoglobin, absolute number of lymphocytes and hemoglobin were found to be significantly decreased. Survival was poor in HD patients with anemia, lymphopenia, thrombocytopenia, hypoalbuminemia and high levels of ALP.

Ref: 77

Abstract No: 39

HODGKIN’S DISEASE IN CHILDREN: DEMOGRAPHIC DATA AND RESULTS OF OUR CLINIC

1Ferhan Akici, 1Gönül Aydogan, 1Zafer Salcioglu, 1Serdar Sander, 1Deniz Tugcu, 1Arzu Akcay, 1Hulya Sen, 1Aysegul Kiyak, 1Huseyin Aldemir, 1Fulya Yaman 2Bakirkoy Women and Children Diseases Education Hospital, Istanbul, Turkey 3Istanbul University Oncology Institute Division of Radiation Department, Istanbul, Turkey

The epidemiologic pattern of Hodgkin’s Disease in developing countries is different when compared with developed countries. In this study conducted between September 1990 and February 2007, the demographic data and results of therapy of 60 patients under the age of 16 years with biopsy-proven Hodgkin’s Disease in our clinic are presented. The median male ratio was 2: 5. 1. The median age was 6. (5-16) years; 73% were younger than 10 years of age. According to the Rye system, 35 cases (58%) were classified as mixed cellularity, 1 (2%) as lymphocyte depleted, 23 (38%) as nodular sclerosis and 12(2%) as lymphocyte predominant type. Four patients (7%) was classified as stage 32(53%) as stage II, 15 (25%) as stage III, 9(15%) as stage IV. Twenty six (42%) patients had B symptoms. Twenty nine (49%) presented with bulky lymph nodes (> 6 cm), 20(33%) with bulky mediastinum. Staging procedures included selective exploratory laparotomy in 10 patients; in 3 of whom, there was a change in the stage. Treatment consisted of two cycles of ABVD chemotherapy for stages I and IIA, four cycles of ABVD for stages IIB and IIIA, six cycles of MOPP/ABV for stages IIIB and IV. All children received involved field radiotherapy of 15 Gy ±5 years old, 20 Gy if 6-10 years old, 25 Gy if 11 years old. An additional 5 Gy was given in patients presenting with bulky mediastinum, bulky lymph nodes and in patients in whom a complete response could not be attained following chemotherapy. Six patients relapsed 8, 9, 11, 22, 36 and 98. months after cessation of therapy respectively and are alive after salvage therapy. Forty-nine of the remaining 60 patients are alive with no evidence of disease, six are lost to follow up and five died due to sepsis or to progression. The 5 year overall survival was 100% for stages I and II, 90% for stages III and 78% for stages IV respectively. In conclusion, the is a predominance of mixed cellularity subtype, male sex and younger age in our population. Results obtained with a combined modality therapy consisting of chemotherapy, modified according to stage, and low dose involved field radiotherapy are satisfactory.
HODGKIN’S LYMPHOMA: A RETROSPECTIVE ANALYSIS OF 44 PATIENTS
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Izmir Training and Research Hospital, Izmir, Turkey

Hodgkin lymphoma, first described by Thomas Hodgkin, is a neoplasm of lymphoid tissue. Morphologic and immunophenotypic features can distinguish four subtypes of classic Hodgkin lymphoma. The incidence is 2.3-3/100,000 per year and there is a bimodal distribution of Hodgkin lymphoma in western countries with two peaks at 15-34 years and over 60 years. The nodular sclerosis subtype predominates in young adults whereas the mixed cellularity subtype is more common in the pediatric population and in older age population. In this study we evaluated retrospectively a total of 44 patients with Hodgkin lymphoma who were followed in our centre from 1995 to 2005. The mean age of patients was 44.2 years (range 24-72). Twenty-nine patients (66%) were male and fifteen patients (34%) were female. According to histopathologic subtype nodular sclerosis was the most common subtype (52%) and lymphocyte depleted subtype was only (7%). Ann-Arbor classification system was used for staging. Seventeen patients (38%) were stage I, fifteen patients (34%) were stage III, seven patients (16%) stage I and five patients (12%) were stage IV. More than 50% of all patients had B symptoms at presentation. B symptoms were seen at thirtyfive patients (80%) with advanced stage and at nine patients (20%) with localized stage. Peripheral lymphadenopathy was determined at the majority of patients and cervical involvement was 64%. Splenomegaly was found in 4 patients (9%). 11% of patients had mediastinal lymph node involvement at presentation and were seen the most common in nodular sclerosis subtype. When laboratory findings examined the mean erythrocyte sedimentation rate was 72.2 mm/h, the mean hemoglobin level was 11.1 g/dl and the mean LDH level was 393.5 U/L at presentation. At the same time there was leucopenia in 16% and leucocytosis in 18% of patients. As a first line therapy 80% of patients received ABVD and 14% received MOPP regimen. Three patients with localized stage received radiotherapy. According to histopathologic subtypes there were no differences between the patients in 5 years survival. Survival rates were 76% in patients with localized stage and 33% in patients with advanced stage. This was statistically significant (p <0.05). As a result most of our patients were in advanced stage at presentation. 52% of patients were nodular sclerosis Hodgkin lymphoma as were seen most common in young adults.

NON-HODGKIN’S LYMPHOMA
Ref. No: 15 Abstract No: 42

CASE REPORT: MANTLE CELL LYMPHOMA WITH PULMONARY INVOLVEMENT AT PRESENTATION
1Serkan Ocakci, 1Nur Akad Soyer, 1Murat Tombuloglu, 2Nazan Ozhan
1Ege University, Department of Internal Medicine, Division of Hematology, Izmir, Turkey 2Ege University, Department of Pathology, Izmir, Turkey

A 50-year-old man presented with 39 degrees celsius fever, dyspnea, fatigue, weight lose and night sweats was referred to our hospital because that he had marked leukocytosis, 74000/mm³. His physical examination revealed multiple cervical lymphadenopathies, a right suprACLavicular lymphadenopathy, massive splenomegaly, prolonged expiration, widespread rhonchi, and bilaterally basal pulmonary crackles. His hemoglobin level was 8. 4 g/dl and platelet count was 145000/mm³. Blood smear showed 80% atypical lymphomononuclear cells. A chest radiograph demonstrated a right pulmonary consolidati-on which was later confirmed as pulmonary nodules and consolidation suggestive of pulmonary lymphoma involvement by chest CT. Patient was diagnosed as mantle cell lymphoma by bone marrow aspiration biopsy and flow cytometry. No clinical or radiological response was seen after empiric antibiotherapy so a bronchoscopy was performed. Bronchoscopy did not show a mass lesion but a transbronchial biopsy showed mantle cell lymphoma involvement. Two cycles of Hyper-CVAD regimen were given. The patient could not tolerate Hyper-CVAD. Six cycles of Rituximab- CHOP regimen were given. Mantle cell lymphoma is an aggressive disease with a poor prognosis. Although systemic involvement frequently occurs in mantle cell lymphoma, primary pulmonary involvement has not been reported so far.
A NON HODGKIN'S LYMPHOMA CASE WITH OVARIAN INVOLVEMENT
Abdullah Hachanefoğlu, Naile Gökkaya, Pınar Tarkun, Emel Gönülül
Kocaeli University, Hematology Department, Kocaeli, Turkey

A 38 years old woman was admitted to our hospital with weakness, loss of appetite, abdominal distention and pain in June 2005. There were symptoms like weakness, loss of appetite, profuse sweating at night, fever, pruritis, abdominal pain and distension at her history. There was no peripheral lymphadenopathy and/or hepatosplenomegaly. In laboratory examination, whole blood count was normal except normocytic anemia (Hb: 11.3 g/dl, HCT: %32). Erythrocyte sedimentation rate was 49 mm/h. In the biochemical values there were no abnormality except elevation of LDH (LDH: 383 IU/L). The viral markers were negative (HBV, HCV, CMV, EBV). The lesion that is 112x87 mm with heterogenous echo and irregular border was found in right adnexial area at pelvic ultrasonography. The thoracal tomography was evaluated as normal. In the abdominal tomography a lesion the 12x10 cm diameter in the left pelvic area was found. There was also lymphadenopathy that is 4x3 cm in diameter with regular border and hypodens in paraaortic area. The patient was operated with diagnoses of over carcinoma by gynecologist and applied salpingo-oophorectomy + right wedge resection + omentectomy. The pathological report of specimen revealed tight cytoplasm, coarse chromatine and angiosentritic arrangement in places at the tumoral cells. In immunohistochemical study CD 45 RO focal (+) ; CD 3 (-) ; CD 5 (-) ; CD 34 (-) ; HLA DR (-) ; MPO (-) and CD 10 (fokal (+) ) was found. These findings made us thinking of B lymphoblastic lymphoma/ leukemia infiltration was shown this time. Ida-FLAG chemotherapy regimen was given with the diagnose of stage IE, B lymphocytic lymphoma. After the end of chemotherapy the patient had her first remission and discharged from the hospital. She was periodically controlled but in June 2006, she addmitted to the emergency room with terrible bone pain. A thrombocytopenia (849000) was seen and the other parameters were normal in complete blood count. Erythrocyte sedimentation rate was 42 mm/hour and LDH was 1230 U/L. In bone marrow examination lymphoblastic lymphoma/ leukemia infiltration was shown this time. Ida-FLAG chemotherapy regimen was given to the patient and at the end of this therapy bone marrow was normal. And she is already is being followed up in our policlinic. DISCUSSION: Primary ovarian lymphoma or secondary ovarian involvement as initial manifestation of lymphoma is rare (1). As in our patient, the most common presenting signs or symptoms of malignant lymphomas involving the ovaries are abdominal or pelvic pain or mass. (1) Malignant lymphoma affecting the ovary can be divided into two types; primary and disseminated. And most patients with ovarian lymphomas are treated with surgery and chemotherapy. (3). Our case was secondary ovarian involvement and we used firstly the Hyper-CVAD and then FLAG-IDA chemotherapy regimens.
ring 3 cm at the liver hilus, surrounding the portal vein and pancreatic head. The histopathological examination of excisional submandibular lymphadenopathy revealed AILD. The stage was IIIA, and CHOP chemotherapy was administered soon. Control CT showed significant reduction in the sizes of the lymphadenopathies, hence AILD was in very good PR. A PET scan was performed in order to detect the possible lymphadenopathies in the abdomen and other regions. His PET scan demonstrated a focally increased FDG at the inferior pole of the left thyroid lobe. Thyroid USG performed, 2 cm and 1 cm hypoechoic view at isthmus and left lobe, respectively. Thyroid fine needle biopsy was performed and revealed suspicious malignancy, and hence a total thyroidectomy performed. The pathologic examination demonstrated papillary carcinoma of the thyroid. There is no clear-cut data for the incidental findings of the secondary cancers. Most data are case reports. As increased number of PET scanning performed, secondary malignancies could be found more frequently. The frequency of thyroid cancer as incidental findings has been reported to be 1% in the published literature. As in our case, we do not have a pretreatment PET scan which prevented us to make a healthy judgement. The another important matter that if a patient PET scan show uptake in thyroid with other regions, it can be related to the primary disease, so secondary malignancies may be overlooked. It is highly probable that most cancer patients die without detecting their secondary malignancies. In conclusion, a positive PET finding in other solid organs in patients with AILD and lymphomas should not only be regarded as the metastasis of the primary malignancy, but also a possibility of a secondary malignancy should be undertaken.

Ref. No: 58 Abstract No: 46

NASAL NK/T CELL LYMPHOMA: CASE REPORT

Arzu Ergen, Barkın Sakallıoğlu, Nergiz Dağoğlu, Yavuz Dizdar, Fulya Yaman Ağaoğlu, Emin Darendeliler
Istanbul University, Medicine Faculty of Istanbul, Department of Radiation Oncology, Istanbul, Turkey

Non-Hodgkin lymphomas originating from nasal cavity, paranasal sinuses and hard plate are a different subgroup of lymphomas. This rare group of lymphomas are characterised with progressive erosive lesion and destruction of bone, cartilage and soft tissue. Our male patient age 52 applied the hospital with sensation of destruction of bone, cartilage and soft tissue. Our male patient age 52 applied the hospital with sensation of burning in his nose and in his physical examination in ENT policlinic an acneiform lesion beginning to ulcerate and rhinorea was found. He was diagnosed as maxillary sinusitis, had aspiration and was prescribed antibiotics but after two weeks of antibiotics treatment he did not recover. Later a biopsy was performed from the nasal cavity and he was diagnosed as extra nodal NK/T cell lymphoma (CD3 (+). CD56 (+), CD 45Ro(+), CD 16(+), CD 7(-), CD20(-) and was referred to oncology policlinic. In his CT scan a soft tissue mass was seen in right maxillary sinus extending in right orbit, skin and extra coanal area with concommitting submucosal soft tissue masses in nasopharynx, oropharynx and hypopharynx. As being diagnosed Stage II NHL, CHOP chemotherapy was planned. In his control during the chemotherapy lesions were found to be progressing both on CT scan and clinical examination, he was referred to our department and was treated with 6 MeV photons with a dose of 180 cGy daily fractions at total dose of 45 Gy RT in one anterior oblique field. No serious acute reactions were observed during radiotherapy and after the treatment lesions were clinically stable. After a month in his control CT scans a new lesion was found in his tonsil, which was in his previous radiation field and because of this new lesion, second line chemotherapy was planned but at that time his condition deteriorated and died before starting his first cycle of chemotherapy. Nasal NK/T cell lymphomas are morphologically, immunophenotypically and genotypically similar to non-nasal extra nodal NK/T cell lymphomas but have an aggressive clinical behaviour. In different clinical trials 3 year overall survival in nasal lymphomas is between %24 and %64. 5 year overall survival in nasal NK/T cell lymphomas is found to be %25 and in non-nasal NK/T cell lymphomas it is %10. Prognosis in NK/T cell lymphomas is poor and more aggressive and effective treatment modalities are needed.

Ref. No: 82 Abstract No: 47

PRIMARY MEDIASTINAL B-CELL NON-HODGKIN’S LYMPHOMA PRESENTED WITH CARDIAC INVOLVEMENT

İnci Alacacioglu, Nurhilal Turgut, Guner Hayri Ozsan, Özden Piskin, Mehmet Ali Ozcan, Fatih Demirkan, Bahri Akdeniz, Mustafa Secil, Omer Kozan, Bulent Ündar
Dokuz Eylül University Faculty of Medicine Department of Hematology, Izmir, Turkey
Dokuz Eylül University Faculty of Medicine Department of Cardiology, Izmir, Turkey
Dokuz Eylül University Faculty of Medicine Department of Radiology, Izmir, Turkey

Primary mediastinal large B-cell lymphoma (PMLBCL) represents a distinct clinical entity with unique clinicopathologic and genetic features. It accounts for 2% of patients with non-Hodgkin’s lymphoma (NHL), is usually limited to the intrathoracic organs, but may spread to visceral organs such as liver, kidneys and the central nervous system. Lymphoma with cardiac involvement is very uncommon and often very difficult to detect while the patient is alive. 64-year-old female patient presented with angina and dyspnea on exertion to emergency room. She was diagnosed as unstable angina pectoris with electrocardiographic and clinical findings. The mass that extended from free wall of right atrium, passing through tricuspid valve, to free wall of right ventricle and constricted the pulmonary valve, another one at left atrium was seen at her transesophageal echocardiography (ECHO). Ejection fraction rate (EFR) was 50%. At her cardiac MRI, the huge mediastinal mass that filled whole mediastinum, encircling main vascular structures, invading cardiac valve and wall, and right pleural effusion were seen. Surgical biopsy by mediastinoscopy revealed the histology of diffuse large B-cell lymphoma. Bone marrow biopsy and abdominal computerized tomography were normal. The patient was put on R-CHOP chemotherapy with near follow-up due to risk of sudden death following rapid tumor regression. Currently, after 2 cycles of chemotherapy all symptoms of the patient disappeared as shown by improved EFR from 50% to 65% on ECHO and re-evaluation by imaging procedures demonstrated good partial remission.
THE ROLE OF RITUXIMAB ON AUTOLOGOUS TRANSPLANTATION FOR NON HODGKIN’S LYMPHOMA

Sinem Civriz Bozdogan, Pervin Topcuoglu, Ender Soydan, Mutlu Arat, Osman Ilhan, Haluk Koc, Meral Bekcsem, Akin Uysal, Hamdi Akan, Onder Aslan, Nahide Konuk, Muhit Ozcan

Ankara University Faculty of Medicine, Ankara, Turkey

High dose chemotherapy and autologus stem cell transplantation is the standard treatment regimen for relapsed non Hodgkin lymphoma (NHL) patients. We aimed to analyze the role of pretransplant rituximab therapy on graft function and outcome of autologous transplantation. After exclusion of T cell lymphoma and lymphoblastic lymphoma patients, we studied on 68 NHL patients and analyzed pretransplant and posttransplant outcomes retrospectively. Median age was 42 years (17-64), 47 male and 21 female patients included to the study. Seventy percent of patients' histopathology was diffuse large B cell lymphoma. Patients were grouped as those who never treated with rituximab before transplantation and those who had rituximab during salvage treatment or mobilization. Clinical features and transplantation outcomes are shown in the table. Rituximab treatment during pretransplantation period has no significant effect on remission ratios. (p>0.05). The outcome after transplantation also is also similar in both groups. The median follow up is 52 months for all patients. Also, no significant effect is observed on 2 years disease free and overall survival with rituximab. These outcomes have to be supported with larger prospective randomized trials.

<table>
<thead>
<tr>
<th>Rituximab (+) (n: 17)</th>
<th>Rituximab (-) (n: 51)</th>
<th>p</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age (median) 47(20-64)</td>
<td>41(17-54)</td>
<td>0.016</td>
</tr>
<tr>
<td>Sex Female</td>
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<td>39</td>
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<tr>
<td>Male</td>
<td>8</td>
<td>13</td>
</tr>
<tr>
<td>Pretransplant response Chemosensitive</td>
<td>12 (70.5%)</td>
<td>30 (58.8%)</td>
</tr>
<tr>
<td>Chemoresistant</td>
<td>5 (29.5%)</td>
<td>17 (41.2%)</td>
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<tr>
<td>Transplant outcome: CR</td>
<td>13 (81.3%)</td>
<td>35 (71.4%)</td>
</tr>
<tr>
<td>PR (relapse posttransplant)</td>
<td>1 (6.3%)</td>
<td>5 (8.2%)</td>
</tr>
<tr>
<td>NR</td>
<td>2 (12.5%)</td>
<td>10 (20.4%)</td>
</tr>
<tr>
<td>Relapse posttransplant</td>
<td>13 (76.5%)</td>
<td>26 (51.3%)</td>
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<tr>
<td>Trans related mortality</td>
<td>5/17</td>
<td>15/36</td>
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<tr>
<td>2 years disease free survival</td>
<td>51.9%±5.6%</td>
<td>46.9%±7.8%</td>
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<tr>
<td>2 years overall survival</td>
<td>56.3%±7.7</td>
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Ref. No: 95 Abstract No: 49

FACS ANALYSIS OF PERIPHERAL T-CELL LYMPHOMA

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Peripheral T-cell lymphoma consists of a diverse group of post-thymic tumors bearing a mature T-cell phenotype and, excluding mycosis fungoides, comprises approximately 10-20% of the non-Hodgkin’s lymphomas in the United States. This category of non-Hodgkin’s lymphomas exhibits considerable morphological, immunological, and clinical diversity and is generally considered to be a high-grade malignancy. Flow cytometry (FC) has become the routine technique in the evaluation of hematopoietic neoplasms. We diagnosed a case of peripheral T-cell lymphoma by using flow cytometry before paraffin-embedded biopsy specimens of lymph nodes. A 32-year-old male presented with fatigue, decreased appetite and multiple lymphadenopathy in his neck. Immunophenotypic analysis of the peripheral blood by flow cytometry demonstrated that the majority of cells were CD4-positive T-cells with a partial loss of CD3 but strongly expressed CD10. After a needle node pulling and cells isolated, similar results was shown by using flow cytometry. Histopathology revealed a diffuse mixed-cell infiltrate of lymphocytes, histiocytes, and numerous eosinophils, which extended throughout the reticular dermis and into the subcutaneous adipose tissue. Scattered lymphocytes have enlarged, hyperchromatic nuclei. Lymphocytes extend into follicular and sebaceous epithelium but spare the overlying epidermis. Most peripheral T-cell lymphoma are aggressive malignant conditions with only cutaneous anaplastic large-cell lymphoma and mycosis fungoides typically displaying indolent clinical courses. This case demonstrates a rare presentation of peripheral T-cell lymphoma. FACS methodology has the advantage of rapid turn-around time as well as high sensitivity, enabling patients with lymphomas. In experienced hands, flow cytometry plays a valuable and complementary role to histology and immunohistochemistry in diagnosing lymphomas.

Ref. No: 100 Abstract No: 50

ROLE OF HEPATITIS B VIRUS AND HEPATITIS C VIRUS INFECTIONS IN CLINICAL OUTCOMES OF NON-HODGKIN LYMPHOMA

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1Shiraz Transplant Research Center, Shiraz, Iran 2Hematology-Oncology Research Center, Shiraz, Iran

Background: Non-Hodgkin lymphoma (NHL) is important types of lymphoproliferative disorders and multiple risk factors have different role in NHL presentation. Infections like, Hepatitis B virus (HBV) and hepatitis C virus (HCV) infections may have an effective role in NHL clinical outcome. Objectives: In this research for determination the role of HBV and HCV infections in NHL patients, the molecular prevalence of these viruses were studied. Material and Methods: In this retrospective and cohort study, 70 and 100 EDTA treated blood samples were collected for 2 years from NHL patients and healthy control group, respectively and the prevalence of HBV and HCV viral genomes were analyzed. Results: HBV and HCV infections were detected in 14% and 20% of NHL patients, respectively. HCV infection was detected in 7% of healthy control group, but HBV infection was not detected in normal control group. HBV and HCV co-infection also was detected in 5.7% of NHL patients. Conclusion: For high prevalence of HBV and HCV infections and co-infection in NHL patients, monitoring of these viral infections may have a role in therapeutic management of NHL patients are needed.
LYMPHOMATOID PAPULOSIS: A CASE REPORT AND TREATMENT RESULTS OF LOCAL THERAPY RESISTANT CASES
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Cerrahpasa Medical Faculty, Istanbul, Turkey

Lymphomatoid papulosis represents a benign, chronic, recurrent, self-healing, papulonodular, and necrotic skin eruption. It is a rare disease; the prevalence is estimated to be 1. 2-1. 9 cases per million population and 10-20% of patients may develop a lymphoid malignancy. We presented three years old boy who had been followed up for 18 months at department of dermatology with complaints of erythematous ulcerative, painful pruritic lesions. Skin punch biopsy revealed lymphomatoid papulosis type A and 60 cures of PUVA light therapy had been applied however no response was observed. The patient was referred to department of hematology and oncology for systemic chemotherapy. There is no curative treatment available for lymphomatoid papulosis and usually managed by observation, intralesional steroid injection, topical bexarotene, imiquimod (Aldara), ultraviolet light therapy, or low-dose methotrexate. Trials with SGN-30 (anti-CD30 mAb) are in progress. We presented this rare case to discuss treatment modalities for local therapy resistant cases.

CLINICAL CHARACTERISTICS AND TREATMENT RESULTS OF PEDIATRIC NON-HODGKIN’S LYMPHOMA
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Purpose: The aim of this study is to evaluate the clinical characteristics of the non-Hodgkin Lymphoma (NHL) patients and treatment results of modified NHL-90 protocol in our clinic. Methods: From January 1996 to February 2007, 54 newly diagnosed children with NHL were enrolled. The Murphy classification was used for staging. The patients were stratified into three groups according to risk factors (stage, LDH, CNS on bone marrow involvement) and treated either with a modified NHL-90 (Berlin-Frankfurt-Münster) protocol. The use of 1 gr/m2 Methotrexate instead of 5 gr/m2/24hr was the only important modification in BFM-90 protocol. Result: Fifty-four children (16 girls, 38 boys) with a median age of 7 years (range 2-15 years) were treated in our clinic. Of these patients, 10(18,5%) had T-cell, 41(76%) had B-cell,3(5,5%) had anaplastic Large Cell Lymphoma. There were 3 patients in stage I, 8 in stage II, 29 in stage III, 14 in stage IV. In 28 patients the primary tumor was in abdomen, in 9 at the head and neck region, in 9 at thorax, and remaining patients had disseminated disease. Complete remission occurred in 43 patients (80%), partial remission in 8 patients (15%) and progressive disease in 1 patient (2%). Only 2 patient died of tumor lysis symptom at prephase. At a median follow up to 54 months (2-135 months) the 5 years overall survival (OS) for all patients was 66%, and event free survival (EFS) was 61%; factors associated with lower EFS by univariate analysis were risk groups, and LDH level (500 IU/L). But there was no statistically significant difference (p=0,90) in EFS. The major toxicity were myelosuppression and mucositis, but these conditions were tolerated and manageable. Conclusion: Intensive, short chemotherapy regimen appears to be superior regimen when compared to others regimens. The treatment results in our clinic are comparable to those of BFM group. This modified NHL-BFM 90 protocol is very effective for children and adolescent with Burkitt Lymphoma and Large Cell Lymphoma.

CLINICOPATHOLOGIC FEATURES AND TREATMENT RESULTS OF NON-HODGKIN’S LYMPHOMAS IN ELDERLY PATIENTS: A RETROSPECTIVE ANALYSIS FROM “DENIZLII LEUKEMIA-LYMPHOMA-MYELOMA STUDY GROUP” (DLLMSG)
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Purpose: The aim of this study is to evaluate the clinicopathologic features and treatment results of non-Hodgkin’s lymphoma (NHL) in the elderly. “Denizli Leukemia-Lymphoma-Myeloma Study Group” (DLLMSG) was nearly established to register the data of lymphoma and leukemia patients in our city in Western Anatolia. So, we have carried out a retrospective analysis of Non-Hodgkin lymphomas (NHL) in elderly patients (age>60) followed at our hematology centers, with the purpose of evaluating the clinicopathologic features and treatment results. Patients and methods: Thirty-one elderly lymphoma patients were assessed with regard to their characteristics including age, gender, histologic distribution, stage, extranodal involvement, presenting symptoms, and also treatment responses. Results: Among 31 elderly patients with NHL, 16 (51%) were male and 15 (49%) were female. The overall median age was 68. 5 years (range; 61-87). Clinical presentation was characterized by superficial lymphadenopathy (70. 9%). According to the Ann Arbor staging system, the vast majority of patients (77. 4%) were advanced stage. The patients were classified according to the World Health Organization (WHO) system. The most commonly observed histopathologic type was Diffuse large B-cell lymphoma (DLBCL) were seen in 16 (51. 6%) patients. International Prognostic Index (IPI) scores were high in 13 patients (42%). Extranodal involvement was found in 8 (25. 8%) patients. Majority of patients were treated with full dose R-CHOP regimen. Drug-dose reduction (25%) were done in 7 patients (22. 6%). Complete remission (CR) was obtained in 19 patients (61. 2%), 6 (16. 1%) of whom relapsed. Grade 3-4 hematologic toxicity was observed in only 16. 1% of cases and there was one treatment-related death because of septic shock. Conclusion: Clinicopathologic features of these patients resembles with younger patients. In addition, full dose R-CHOP regimen were effective and relatively tolerable as well as in younger adults.
ARG: A POTENTIAL BIOMARKER FOR DLBCL STAGING

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ARG is a proto-oncogene and a member of a tyrosine kinase proteins family. It has great importance in many cancers, but there is no conclusive evidence on its role in DLBCL staging. The aim of this study was to evaluate importance of ARG, in cancer's staging. Sixteen DLBCL and 4 reactive lymph nodes (as control group) samples, after staging using Ann Arbor staging system, were used in this study. Formaldehyde fixed paraffin embedded blocks were prepared from the samples. After sectioning the samples were hybridized with fluorescently labeled probes against ARG; FITC labeled and GAPDH; Rodamine labeled (as control house keeping gene). After capturing the pictures using CCD camera, the intensity of green and red colures were measured and ratio between green/red, that demonstrate changes in ARG expression, were calculated. The mean ratio of green/red (ARG expression) was significantly different between reactive lymph node, stage I, II and III of DLBCL. ARG expression was different between all groups except for between stage III and VI of DLBCL. The observed changes in ARG expression is a potential biomarker for DLBCL staging. In addition specific inhibitors of ARG can be considered as new chemotherapy agents in DLBCL treatment.

A CASE OF BURKITT'S LYMPHOMA WITH NECROTIZING GRANULOMATOUS REACTION

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Introduction: Burkitt lymphoma is a aggressive, heterogenous B cell lymphoma. Burkitt lymphoma related to Epstein Barr Virus and usually seen in children. Although well recognized in T-cell NHL and Hodgkin’s disease, Burkitt lymphoma with granulomatous reaction has been rarely reported in B cell lymphomas. Case Report: An 6-year-old boy presented with lymphadenopathy in head and neck region. The ultrasound and computed tomography procedures revealed a mediastinal mass. A surgical sample from the servical lymph node showed morphologic and immunophenotypic features of Burkitt’s lymphoma with large necrosis and granulomatous reaction. In the microscopical examination there was a diffuse infiltrate of atypical lymphoid cells with numerous mitoses and prominent starry-sky pattern because of the presence of multiple tingible body macrophages. Prominent infiltration of epitheloid histiocytes, forming small clusters and granulomas of different size were remarkable. Multinucleated giant cells were identified within granulomas. Discussion: In this study, we present the a case of sporadic Burkitt lymphoma with an extensive epitheloid cell granulomatous reaction. Burkitt’s lymphoma with granulomatous reaction has been rarely reported in B cell lymphomas. A granulomatous reaction may also be caused by a concomitant infection by m. tuberculosis, yeast, fungi or other microorganisms.

LYMPHOMA EXPERIENCE OF LAKES DISTRICT FROM SÜLEYMAN DEMIREL UNIVERSITY SCHOOL OF MEDICINE

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Purpose: Lymphoma is one of the most common cancers in adult patients. Especially Non-Hodgkin’s lymphoma incidence is increasing. Neither national nor regional cancer statistic was not reliable in Turkey. Method: We analyzed principle epidemiologic data of patients with malignant lymphoma in Lakes District from Turkey. All patients were adopted from SDU cancer registry data. We analyzed 154 lymphoma patients. Of 154 patients 27 were diagnosed before 2002 when the department of hematology and medical oncology was not established in SDU. Results: We analyzed 154 lymphoma patients of whom 44 were Hodgkin’s (HL), 110 were non-Hodgkin’s lymphoma (NHL) patients. Median age was 64.5 (22-82) years for NHL. Sixty patients were male and fifty were female. We had 15 extranodal lymphoma patients. One had multiple extranodal (parotis, terminal ileum, pancreas), 7 gastric, 4 central nervous system, 2 skin, 2 parotid involvement. Sixty percent of the cases diagnosed as diffuse large B cell lymphoma. Most of the patients were at advanced stage when diagnosed, as stage III 29.9% and stage IV 41%. Most of the cases were put on CHOP (38%) and R-CHOP (48%) protocols as up front treatment. Median age was 46 (18-89) years for HL. Thirty patients were male and 14 were female. Rate of early stage and advanced stage cases were as follows: Stage I+II: 48.5%, stage III+IV: 51.5%. Most of the patients were diagnosed as mixed cellularity HL 59.4%, nodular sclerosis HL 31.2%. Most of the patients were treated with ABVD chemotherapy protocol (88.2%). Conclusion: Any effort done to realize cancer status of Turkish cancer population would add more progress to understand the evaluation of cancer in Turkey.

UTILITY OF PERIPHERAL BLOOD FLOW CYTOMETRY TO INVESTIGATE THE PERIPHERALIZATION OF B-CELL MALIGNANT LYMPHOMAS

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Aim: To determine the diagnostic value of investigating B-cell clonality by flow cytometric analysis of peripheral blood samples in B-cell lymphomas. Lymphomas are known to have a relatively high rate of peripheral blood involvement as well as lymph node areas, bone marrow and several organs. Methods and results: Blood flow immunophenotyping studies for B-cell clonality were performed in 43 cases who had persistent enlargement of lymph nodes or spleen and 11 healthy adults. The diagnosis included B-cell NHL in 29 cases, T-cell NHL in 1 case, Hodgkin’s disease in 2 cases and metastasis from solid cancers in two cases. The remaining 7 patients had a variety of nonmalignant diseases. Flow cytometric immunop-
hemotyping of an EDTA, anticoagulated peripheral blood was performed in each case, with CD19- PerCp/kappa-fluorescein isothiocyanate (FITC)/lambda-phycocerythrin (PE) (Becton Dickinson, Franklin Lakes, NJ, USA). B-cell clonality was determined according to five different patterns: 1. Abnormal kappa/lambda ratio (>3) or lambda/kappa ratio (>2). 2. Abnormal B-lymphocyte subpopulation with different CD19 expression. 3. Abnormal B-lymphocyte subpopulation positive for kappa and lambda on the same cell population. 4. Abnormal B-lymphocyte subpopulation negative for both kappa and lambda on the same cell population. 5. Abnormal CD19 expression(increased or decreased) on B-lymphocytes. We found that 18/29(62%) of patients with B-cell NHL had abnormal findings consistent with B-cell clonality in peripheral blood: 16/29 (55%) had abnormal kappa/lambda ratio, the kappa light chain being dominant in 14; 15/29 (52%) had abnormal CD19 expression, increased CD19 expression being common in 12; 7/29 (24%) had abnormal B-lymphocyte subpopulation negative for both kappa and lambda on the same cell population; 1/29 (3%) had abnormal B-lymphocyte subpopulation positive for both kappa and lambda on the same cell population, and finally 2/29 (7%) had abnormal B-lymphocyte subpopulation with different CD19 expression. Kappa/lambda ratio was also higher in patients with B-cell NHL than the patients with other causes of enlarged lymph nodes and spleen (p<0.001). Interestingly, B-cell clonality was detected by peripheral blood flow cytometric immunophenotyping in three patients who presented with isolated splenomegaly and later diagnosed as B-cell NHL.

Conclusions: We concluded that a peripheral blood flow cytometric immunophenotyping study could be used to investigate the peripheralization of B-cell lymphomas and abnormal kappa/lambda ratio discriminates B-cell lymphomas from other causes. Investigation of B-cell clonality by peripheral blood flow cytometry is also valuable for the differential diagnosis of splenomegaly in cases of suspected B-cell lymphoma.

**LYMPHOPROLIFERATIVE DISORDERS**

Ref. No: 110 Abstract No: 59

OPPORTUNISTIC INFECTIONS IN CASES WITH HAIRY CELL LEUKEMIA

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Hairy cell leukemia is a B-cell disease that the abnor-
mal cell has prominent cytoplasmic projections and contains the tartrate-resistant isozyme 5 of acid phosphatase. This cell infiltrates bone marrow, liver and spleen, result-
ing in organomegaly and pancytopenia. A fewer may be due to the disease as well as infections. Hairy cell leukemia is associated with gram-positive and gram-negative bacte-
rial, atypical mycobacterial and invasive fungal infections. Infection is the primary cause of morbidity and mortality in this disease. In this report we present 7 cases dealt
with in the hematology department of Baskent University, from 2004 to 2007. Two of the patients were female and five were male. Average age was 45 years. Two patients
were given the interferon therapy as an initial treatment, but they did not tolerate treatment well. One of the pa-
tients were performed splenectomy. All patients were given cladribine 0.1 mg/kg continual infusion for 7 days. In 6 patients, fever episodes were observed. In 3 patients pulmonary infection, in 1 patient tuberculous peritonitis and in 1 patient CMV infection was observed. Pulmonary infections that were seen in 3 patients developed before cladribine therapy. CMV infection was seen a month after and tuberculous peritonitis was seen two month after the therapy. There was a correlation between presence of pre-treatment infection and absolute neutrophil count at time of diagnosis. We observed pneumocystis carinii pne-
monia in the patient with the least absolute neutrophil count (120). Pseudomonas aeruginosa and staphylococcus aureus growth was seen on sputum cultures obtained in different times. No febrile episode was seen in the 3 cases with the highest absolute neutrophil count (500-1410) at the time of diagnosis. In 1 patient, pulmonary infection was seen but the case could be easily controlled. In two patients with high absolute leukocyte count at the time of diagnosis, we found CMV infection and tuberculous perit-
onitis after the treatment process. These opportunistic

**EVALUATION THE RESPONSE RATE OF IEV REGIMEN AS SALVAGE THERAPY FOR RELAPSED / REFRACTORY NON-HODGKIN’S LYMPHOMA PATIENTS**

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Aims: Despite advances in the management of agg-
ressive non-Hodgkin’s lymphoma, the treatment of relapsed and primary refractory disease remains a major challenge. Therapy for relapsed/refractory lymphomas should be based only on drugs not included in the front-
line chemotherapy regimens. High-dose chemotherapy or radio-chemotherapy with and without autologus stem cell transplantation is a potentially curative treatment approach. Design and Methods:10 patients with relapsed or refractory non-Hodgkin’s lymphomas received ifosfami-
de & Mesna 2g/m2 daily for 3 days in combination with epirubicin 100 mg/m2. Day 1 and etoposide 150mg/m2 days 1-3. Of the 10 patients with non-Hodgkin’s lymphomas in this study 2 had primarily refractory disease, 8 had developed relapse following primary treatment in less than 6 months. Results: The overall response rate was 90%; it was 50% complete response and 40% partial response. Two proceeded to autologous bone marrow transplantati-
on. Eight patients remain alive in continuous remission with a follow-up of 3-21 months. We treated 10 patients and observed NIH hematoxicity grade 1 neutropenia (10%) and 4 (40%) grade 1 thrombocytopenia (20%), grade 2 anemia (40%) nausea (100%),fever in 80%,neutropenic fever in 20%,UTI in 30%,pneumonia in 20%,of patients, but improved over the three courses of treatment. There was no major toxicity. Further trials of this regimen in this clinical situation are indicated. Tolerance to the regimen was good. The probabilities of overall survival, and the disease-free survival at end of study were 80%, with dura-
tion 13 & 5 months. Additional follow-up is necessary to determine if this improvement in the complete remission rate will confer an increase in the overall survival. Interp-
etation and Conclusions: Our results indicate the efficacy of the IEV regimen in inducing a good remission rate. IEV is a predictable and highly effective in relapsed/refractory patients with aggressive NHL. Key words: IEV regimen, relapse/refractory NHL lymphoma, salvage therapy.
infections were thought to be related to lympholytic effect of cladribine. In hairy cell leukemia, infections can be categorized into 2 as bacterial infections and opportunistic infections that are related to impaired cellular immunity. Infections, especially pulmonary infections, are frequently seen in patients receiving cladribine therapy (totally 28%, severe infection 6%). A correlation between low granulocyte count and bacterial infection incidence was reported in previous studies. However, no relation was found between infection and post treatment lymphopenia. Our results are compatible with the literature.

**CHRONIC LYMPHOCYTIC LEUKEMIA**

Ref. No: 129 Abstract No: 61

**CD4- CD8+ T-CELL PROLYMPHOCYTIC LEUKEMIA: A REPORT OF TWO CASES**

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T cell-prolymphocytic leukemia (T-PLL) is rare neoplasm usually shows CD4+CD8- phenotype. A new variant of T-PLL showing CD-CD8+ phenotype, a lack of stoplasmic azurophilic granules and NK antigens, nuclear polymorphism, and aggressive clinical course, and death within 20 months was reported for the first time in 1987. We reported here two cases with CD8+ T-PLL presenting with dizziness and weakness, respectively. On the basis of results of morphologic examination and flow cytometric analysis of the peripheral blood, a diagnosis of T-PLL was made (Table). One of the patients had a history of splenectomy 20 year earlier than admission. The patients followed up without therapy for three and six months, respectively. The patients showed no progression of their disease while they were under close observation. Our patients were positive for CD7 and negative for HTLV-1 which was ruled out the possibility, adult T-cell leukemia/lymphoma, the lack of stoplasmic granules and NK markers excluded T-cell large granular leukemia, the absence of dermatological findings and positive markers for CD7 and CD8 excluded a diagnosis of Sezary syndrome. Hairy cell leukemia ruled out by flow cytometry. We conclude that it may be difficult to distinguish T-PLL from other lymphoproliferative disease. They may not have always aggressive clinical course

**MULTIPLE MYELOMA**

Ref. No: 10 Abstract No: 62

**A PROTEAZOM INHIBITOR IN THE TREATMENT OF MULTIPLE MYELOMA: BORTEZOMIB**

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Multiple myeloma (MM) is a disease characterised by the proliferation of malignant plasma cells in bone marrow. Although alkylating agents, corticosteroids, antracyclins, vinca alkaloids and thalidomide are used in its treatment, the disease recurs and becomes refractory to treatment. It has been seen that anormal NF-kB signalisation is involved in the pathogenesis of MM and established that this may be inhibited by proteazome inhibitors. Bortezomib is new agent used in MM treatment with this aim. In our clinic, bortezomib was administered to 12 MM cases. The aim of this retrospective evaluation is to evalu-
ate the side effects of bortezomib treatment and establish its efficacy. Mean age of 12 patients (8 male, 4 female) was 60(49-69). Bortezomib was administered as second line treatment in 5 cases, 3rd line treatment in 6 cases and 4th line treatment in 1 case. Bortezomib was administered as single agent to all cases. The dose schedule was planned as follows: maximum 6 courses every 28 days, in each course 1. 3mg/m2 was administered at 1, 4, 8, and 11th days. One case died with the presentation of respiratory failure after the second dose and another case died suddenly with an unknown reason after the second course of treatment. Grade 1 neurotoxicity developed in one case, grade 2 in three cases and grade 4 in one case. In grade 1 neurotoxicity, treatment was maintained at the same dose, in grade 4 neurotoxicity treatment was discontinued. In grade 2 toxicity, dose was reduced to 1mg/m2. In one case, allergic reaction and subsequently rash developed in association with treatment and drug was discontinued. Hematological toxicity was observed in no patient. The responses of the cases to treatment were evaluated using the criteria suggested by SWOG and Blade et al. 1 case did not respond to treatment. Minor response was obtained in 2 cases, partial response in 1 case and clinical remission in 2 cases and complete response in two cases. In conclusion, it has been established that bortezomib is an efficient agent in the treatment of refractory and relapsing multiple myeloma despite its neurological side effects. However, further studies are required with larger patient populations in order to evaluate the efficacy and side effect profile of bortezomib in MM.

Ref: No: 11 Abstract No: 63

GENETIC ABNORMALITIES IN MULTIPLE MYELOMA, THEIR PREVALENCE AND RELATION WITH OTHER RISK FACTORS

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Multiple myeloma is a disease characterised by the proliferation of malignant plasma cells in bone marrow. Although it always has a similar clinical presentation, prognosis may vary considerably. Upon the development of new treatment approaches in the management of MM, prognostic parameters used to determine treatment options have become inadequate and are influenced from other clinical pathologies accompanying MM. Therefore, it was thought that further prognostic markers are required such as cytogenetic characteristics. The aim of the present study is to determine the frequency and types of cytogenetic abnormalities in MM cases and their relation with other risk factors. 50 MM cases undergoing treatment in Ankara Numune Educational and Research hospital were included in the study. In addition to clinical and biochemical evaluation, CRP and Beta 2-microglobulin values were measured. Durie-Salmon and ISS stages were determined and cytogenetic evaluation made. In the conventional cytogenetic evaluation of 42 cases, the most frequently observed anomaly was hypodiploidy 52. 4% (22/42), to be followed by in decreasing order of frequency: near tetraploidy 7. 1% (3/42), hyperdiploidy 2. 4% (1/42), tetraploidy 2. 4% (1/42), del 13q 2. 4% (1/42) and complex karyotype 2. 4% (1/42). In 19% of cases (8/42), other chromosomal abnormalities were observed as well: Namely, ctbh (7q)[10], -22, -Y, del (17)[p13], del (3)[q25]. With FISH evaluation of 48 cases, the most frequently observed anomaly was del 13q, 37.5% (18/48). In the 29 cases evaluated with FISH; t(11;14) was observed at the rate of 24%. 1% (7/29), del 17p 10. 3% (3/29), t(4;14) 3. 4% (1/29), trisomia 11 3. 4% (1/29), trisomia 17 3. 4% (1/29) and CCND1 amplification 6.9% (2/29). It was established that as DS stage increased, DS renal stage and ISS stage increased as well. Since the rate of cytogenetic abnormalities increased as Beta 2-microglobuline level increased, it should be used along with cytogenetic parameters in the determination of prognosis. No relation was found between cytogenetic characteristics (with FISH del 13q, del 17p, t(4;14), t(11;14), DS stage, DS renal stage and ISS stage, which indicates that cytogenetic characteristics is an independent prognostic factor in MM. High expression of del 13q with FISH and complex caryotype anomaly was found to be associated with unfavorable prognosis. It was also seen that in FISH, 17p and t(11;14) rates increased together with del 13q. In order to evaluate the impact of established cytogenetic characteristics on prognosis, we aim to follow the cases for at least five years.

Ref: No: 17 Abstract No: 64

TC-99M MIBI OR F-18 FDG IMAGING?: A COMPARATIVE STUDY FOR EVALUATING PATIENTS WITH MULTIPLE MYELOMA

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Background: Both F-18 fluoro-deoxyglucose-PET (FDG-PET) and Technetium-99m sestamibi (MIBI) scans have been reported to identify sites of disease in patient with multiple myeloma (MM), however their relative utility has not been compared outside a few report. Therefore, the purpose of this study was to compare the diagnostic abilities of the MIBI scan and the FDG-PET scan in the evaluation of MM. Materials and methods: A total of 21 patients with MM (mean age: 61 ±2. 4 years; 7 females, 14 males) were included in the study. Of the 21 patients, 15 were newly diagnosed with previously untreated MM and 6 had relapsed disease after therapy, which was determined by X-ray skeletal survey and hematological/biochemical parameters including complete blood count, liver and kidney function test, erythrocyte sedimentation rate (ESR), serum immunoglobulins, urine light chain excretion, C-reactive protein, β2-microglobulin, and bone marrow plasma cell infiltration. None of the patients with relapsed disease had undergone chemotherapy or radiotherapy during the 6 months preceding the study. F-18 FDG imaging was performed 1 h following administration of 370 MBq of F-18 FDG using a dual head coincidence mode gamma camera. Whole-body MIBI scans were obtained 20 min following iv. administration of 760 MBq of Tc-99m MIBI. Results: There was a positive correlation between the percentage marrow involvement and the number of sites detected on MIBI (P < 0. 001). No such correlation was seen with the number of sites detected on FDG imaging. In 11 of 21 cases (52%), F-18 FDG scan identified addition known active disease at other sites. Ten cases showed unexpected additional sites in patients thought to have limited/stable disease. In 14 of 21 cases (66%), Tc-99m MIBI identified additional sites of disease not seen on routine skeletal survey. Six of 14 cases had known active disease at other sites. Eight cases showed unexpected additional sites in patients thought to have limited/stable disease. Conclusion: F-18 FDG
imaging and Tc-99m MIBI scintigraphy are useful diagnostic tools for detecting otherwise occult sites of involvement by myeloma. However, MIBI imaging can detect more lesions than the FDG scan in patients with MM. The use of MIBI ± FDG imaging should particularly be considered in the evaluation of a patient with presumed limited disease, such as a solitary plasmacytoma, to exclude the presence of other disease sites.

Ref. No: 24 Abstract No: 65
BORTEZOMIB AND DEXAMETHASONE INDUCED TUMOR LYSION SYNDROME IN A CASE OF PLASMA CELL LEUKEMIA
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Tumor lysis syndrome (TLS) is a treatment complication which can be life threatening. This syndrome has been reported more commonly in bulky, hyperproliferative malignancies than solid tumors. Because turnover rate of malignant B cells is low, TLS is seen rarely in plasma cell malignancies. Sensitivity to proteosome inhibitors has been demonstrated in a number of malignancies, particularly multiple myeloma. Bortezomib is the first proteasome inhibitor which has been used as second and third line therapy for patients with relapsed or refractory multiple myeloma. Additionally it has been reported as efficient agent for plasma cell leukemia (PCL). We describe the case of a patient with plasma cell leukemia treated with bortezomib and dexamethasone and developed TLS. A 60 years old man presented increase in lymphoplasmocytoid cells in peripheral blood and bone marrow. He was diagnosed PCL and had been given 6 course of Hyper CVAD therapy. He was in remission after 3th cycle and completed 6 course of chemotherapy. After 3 months, increase of lymphoplasmocytoid cells detected in peripheral blood smear. Bone marrow aspiration and biopsy showed infiltration of lymphocyte, lymphoplasmocytoid cells and plasmoblasts. We decided to give him bortezomib 1.3 mg/m2 i.v on days 1, 4, 8, 11 for three cycles. When starting the first cycle of bortezomib, his thrombocyte count was 55 000/mm3. He tolerated therapy well except for thrombocytopenia. Because of thrombocytopenia we decreased the dose of bortezomib to 1 mg/m2. Dexamethasone 40 mg/day i.v was added on days 1, 2, 3, 4 to the second cycle, leading acute biochemical changes indicative of tumor lysis syndrome, acute renal failure and disseminated intravascular coagulation were seen. His leukocyte count decreased from 4000/mm3 to 1500/mm3 too. After 3 course of hemodialysis he recovered. He had partial response to therapy. We gave only dexamethasone on 1, 2, 3, 4 days as the third cycle because of severe thrombocytopenia and grade 4 neuropathy. TLS has been reported in cases who recieved talidomide, dexamethasone and bortezomib. Only one PCL case who recieved bortezomib has been described with TLS. Because PCL is more rapidly prolifeative disease than multiple myeloma, TLS may be seen in this patients with bortezomib and dexamethasone. Therefore this complication should be looked for during treatment.

Ref. No: 38 Abstract No: 66
SUCCESSFUL TREATMENT OF EARLY RELAPSE OF OCULAR MYELOMA WITH BORTEZOMIB AND STEROID AFTER AUTOLOGOUS STEM CELL TRANSPLANTATION
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Autologous stem cell transplantation (ASCT) can prolong remission duration, overall and progression free-survival in multiple myeloma (MM). Extramedullary plasmacytomas are rare plasma cell tumours originating mostly from the upper respiratory tract and oropharynx. Ocular relapse is rare in MM. Here we present a patient with only ocular relapsed and without evidence of bone marrow progression after ASCT. Ig A kappa myeloma, stage IIIA was diagnosed to the patient a 53-year-old man, according to Kyle-Greipp and Durie Salmon. He was treated with three courses of VAD (vincristine, adriamycin and dexamethasone) therapy. Then he received high dose melphalan (200 mg/m2), followed by the ASCT. After two months ASCT, he had bilateral blurry vision, pain, redness in both eyes and diplopia. We detected 5 mm of right-sided proptosis by Hertel exophthalmometry (base 110, 20 mm right eye, 15 mm left eye). Ocular motility of oculus dexter (OD) was restricted in up and lateral gaze. He has diplopia in up gaze. His color vision was 7 of 12 in the right eye and 10 of 12 in the left eye with Ishihara plates. Best corrected visual acuity was 6/10 in the right eye and 7/10 in the left eye. Intraocular pressures were 19 mmHg for OD and 18 mmHg for oculus sinister. Slit lamp biomicroscopy revealed subconjunctival hemorrhages superiorly and temporally in the right eye and bilateral conjunctial hyperemia with chemosis. Fundus examination of both eyes were unremarkable. Computed tomography and magnetic resonance imaging of orbital revealed a right intraorbital extracanal soft tissue density mass that involved the lacrimal gland and lateral rectus muscle. Prednisolon 1mg/kg/day and bortezomib 1.3 mg/m2 (1, 4, 8, 11 days) were started to the patient. Eye findings were recovered after one month. Ocular relapse should be considered if there are ocular findings. Bortezomib and steroid may be useful for ocular extramedullary relapse.

Ref. No: 44 Abstract No: 67
COMBINED THERAPY WITH BORTEZOMIB AND DEXAMETHASONE IN PATIENTS WITH RELAPSING MULTIPLE MYELOMA
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Objective: To evaluate the efficacy of bortezomib + dexamethasone combined therapy in patients presenting with relapses after treatment of multiple myeloma. METHODS: Patients with relapsing multiple myeloma who were given bortezomib + dexamethasone between the years 2005-2007 were studied retrospectively. Results: A total of 7 patients were assessed in this study (3 males, 4 females). Mean age of the patients was 59.
1 (54-65 years). All 7 patients included in the study were found to be of stage 3 according to the Salmon-Durie staging. Patients had previously been prescribed VAD chemotherapy and complete remission had been obtained, however relapses developed in these patients after a mean period of 6.7 (3-10) months. Four cycles of i. v. bortezomib 1.3 mg/m² (Day 1, Day 4, Day 8, and Day 11) plus p. o. dexamethasone 20 mg/day (Day 1-2, Day 4-5, Day 8-9, Day 11-12) was administered to patients with relapses. Patients were re-evaluated after the 4 cycles. Bone marrow aspiration, serum protein electrophoresis, levels of serum immune globulin, and fixation study in 24-hour-urine specimen were obtained from the patients. Examination results showed that complete remission was obtained in 4 subjects (57.1%), and partial remission was obtained in 2 subjects (28.5%). Another patient however (14.2%), did not respond to treatment, and thus was switched to thalidomide + dexamethasone therapy. This patient went ex 2 months after the onset of thalidomide + dexamethasone therapy. Therapy was continued for 4 more cycles in patients with complete and partial remission and a total of 8 courses of bortezomib + dexamethasone was administered. Patients were re-evaluated after 8-cycles long therapy. The evaluation results showed that complete remission persisted in 4 patients, and that 2 patients with partial remission were still in partial remission after the administration of 8 cycles of therapy. Thalidomide + dexamethasone treatment protocol was initiated in these patients with partial remission. Patients with complete remission have been followed up for a mean period of 12 (11-13) months without any problems arising.

Table 1: Patient Demographics

<table>
<thead>
<tr>
<th>Sex</th>
<th>Age</th>
<th>Duration of relapse (VAD months)</th>
<th>VAD chemotherapy</th>
<th>Complete remission</th>
<th>Partial remission</th>
<th>Unresponsive to therapy</th>
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<tbody>
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<td>Male</td>
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Table 2: Levels of serum IgG and gamma band in protein electrophoresis

<table>
<thead>
<tr>
<th>Levels of serum IgG mg/dl</th>
<th>Before treatment</th>
<th>After treatment with 4 courses of bortezomib</th>
<th>After treatment with 8 courses of bortezomib</th>
<th>Gamma band in serum protein electrophoresis %</th>
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</table>

Ref. No: 66 Abstract No: 68

BORTEZOMIB EFFICIENCY IN MULTIPLE MYELOMA

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Bortezomib is the first proteasome inhibitor to be used in clinical practice. It has shown significant activity in trials of patients with relapsed or refractory multiple myeloma. In the SUMMIT trial, response rate is 27% in relapsed or refractory myeloma patients. The CREST trial showed similar response rate. With the addition of dexamethasone, the overall response rate is higher than monotherapy alone. Bortezomib at dose of 1.3 mg/m² was administered as an injection for eight 3-week cycles. Its adverse events including, thrombocytopenia and peripheral neuropathy. We used bortezomib alone or in combination with other agents. Nine multiple myeloma patients were treated. Bortezomib 1.0 or 1.3 mg/m² was administered days 1, 4, 8 and 11 every 21 days for up to 8 cycles and dexamethasone (40mg orally) on days 1 through 4 with thalidomide (200 mg orally every day). The patients had a median age of 63 years (range, 46-80). The median number of previous treatment was 3 (2-4). Regimens included bortezomib only in 2 patients, bortezomib plus a thalidomide and dexamethasone in 7 patients. Four patients stopped therapy because of adverse events (neuropathy 3; urticarial skin lesions 1). The analysis of patient response to therapy revealed a complete response and or near complete response in 2 patients. We have seen partial response in 2 patients and none response in one patient. Our observation may support that treatment of multiple myeloma patients seems to be intolerant bortezomib used in combination with thalidomide.

Ref. No: 67 Abstract No: 69

RETROSPECTIVE ANALYSIS FOR DEMOGRAPHIC FEATURES OF MULTIPLE MYELOMA PATIENTS, AKDENIZ UNIVERSITY EXPERIENCE

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Key words: Multiple myeloma, demographic features

Introduction: The aim of this study is the retrospective determination of demographical data at the time of diagnosis, first referral clinical characteristics and risk factors for 134 Multiple Myeloma patients with available data records who had been diagnosed, followed and treated at Akdeniz University School of Medicine. Patients and Method: 134 Multiple myeloma patients with available data records, who had been diagnosed based on clinical and laboratory findings, bone marrow
aspiration and biopsy, and radiological examinations of skeleton system, at Akdeniz University School of Medicine Department of Internal Medicine and Department of Haematology, between January 1994 and July 2006 were included into this study. Examinations were carried out retrospectively, based on formal patient files and followup files of Department of Haematology. Findings: Seventy-eight of the patients were male (58%) and 56 of them were female (%42). Male/female ratio was 1,4. Ages of the patients varied between 19 and 90, while the mean age was determined as 60. The highest percent of cases were at the 7th decade (40%). Considering the occupational distribution of the patients, most of them were housewives, officers and farmers (36%, 19%, 16% respectively). Most of the patients referred from town centre (%49). According to body mass index, 53% of patients were over-weight (35%) or obese (18%), 47% of patients had normal weight. The risk factors were, greenhouses for 15%, agricultural disinfection for 13%, story of hair dye use for 9%, history of contact with products of petrol for 1%. The first referral clinics of the patients were, internal medicine for 61 (43%) patients, haematology for 23 (18%) patients, physical medicine and rehabilitation for 9 (7%) patients, orthopedics for 8 (6%) patients, nephrology for 8 (6%) patients, neurosurgery for 7 (5%) patients, chest clinics for 3 (2%) patients and thoracic surgery for 2 (1%) patients. Conclusion: This is a retrospective analysis of data from a single center. With data and findings from our study, we hope to gather data from other centers in order to form Turkiye data at near future.

Ref. No: 68 Abstract No: 70
THE EFFECTS OF PLASMA EXCHANGE ON COAGULATION PARAMETERS, AND PLATELET FUNCTIONS IN PATIENTS WITH MULTIPLE MYELOMA
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Aim: The aim of this study is to investigate the effects of plasma exchange on coagulation parameters, and platelet functions in multiple myeloma patients. Materials (Patients) and Methods: Between February 2004 – July 2006, 10 multiple myeloma patients (male= 6, female= 4) who underwent plasma exchange were included in this study. Coagulation parameters; PT, aPTT, TT, Fibrinogen, AT III, D – Dimer, vWF, Lupus Anticoagulant, Protein C, Protein S, APC Resistance were studied. To determine the platelet functions; Collogen – ADP, Collogen – Epinephrin were performed. Replacement fluids were fresh frozen plasma (FFP), and albumin. Results: FFP was used in 6 patients (60 %), 20 seans (62%). Albumin was used in 4 patients (40%), 12 seans (38%). Significant decrease was detected in D – Dimer levels (p= 0,043) in the sera of the patients who underwent plasma exchange with FFP. After plasma exchange; vWF Factor levels decreased significantly (pre-PE: 97,4±25,06/post-PE: 87,6±22,33, p= 0,001), but there was no changes in APC resistance (pre-PE: 2,11±0,49/post-PE: 2,11±0,52, p= 0,256), and Lupus Anticoagulant (pre-PE: 1,04±0,12/post-PE: 0,99±0,13). There was a significant increase in bleeding time with Collogen – ADP (pre-PE: 174,7±72,77/post-PE: 218,5±76,39, p= 0,047), but there was not a significant change in bleeding time with Collogen – Epinephrin (pre-PE: 199,5±74,81/post-PE: 207,7±60,7, p= 0,603).

During 32 seans plasma exchange procedures, urtiacial symptoms(n= 12, 37, 5 %); itching, flushing, fever etc. were detected. Hypotension (n= 6, 18, 7%), different arrhythmias (n= 1, 3 %), anaphylactic reactions (n= 2, 6 %), and other complications (paresthesia, shivering, dyspnea etc.) (n= 11, 34, 3%) were observed. There was no serious complication, and there was no plasma exchange related mortality. Conclusion: Plasma exchange lowers D – Dimer levels, vWF Factor levels but there is no significant effects on other coagulation parameters. Significant increase in the bleeding time with Collogen – ADP was detected after plasma exchange. Plasmexchange can be performed safely, and complications can be avoided when plasma components were used as replacement fluids in patients with multiple myeloma. Key Words: Multiple myeloma, plasma exchange, coagulation parameters, platelet functions.

Ref. No: 69 Abstract No: 71
RETROSPECTIVE ANALYSIS FOR CLINICAL AND LABORATORY FINDINGS OF MULTIPLE MYELOMA PATIENTS, AKDENIZ UNIVERSITY EXPERIENCE
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Key words: Multiple myeloma, clinical and laboratory findings. Introduction: The aim of this study is the retrospective determination of laboratory data at the time of diagnosis, types of immunoglobulins, first referral symptoms of the patients, stages of disease according to Durie-Salmon and International staging systems for 134 Multiple Myeloma patients with available data records, who had been diagnosed, followed and treated at Akdeniz University School of Medicine. Patients and Method: 134 Multiple Myeloma patients with available data records, who had been diagnosed based on clinical and laboratory findings, bone marrow aspiration and biopsy, and radiological examinations of skeleton system, at Akdeniz University School of Medicine Department of Internal Medicine and Department of Haematology, between January 1994 and July 2006 were included into this study. Examinations were carried out retrospectively, based on formal patient files and followup files of Department of Haematology. Findings: First referral complaints of patients were mostly skeletal pain (waist, dorsum and extremity) and complaints concerning anemia (weakness, fatigue, palpitation, dyspnea) (55% and 31%, respectively). At the time of diagnosis osteoporosis and compression fractures were the most common findings. At the time of diagnosis, 48% of patients had common lytic bone lesions and %31 had no lytic bone lesions. The mean serum calcium, creatinine and haemoglobin levels of patients were 9,8 mg/dl, 2,3 mg/dl and 9 g/dl respectively. When types of immunoglobulins for our cases were examined, it was determined that 59% had IgG, 22% had IgA, 15% had light chain (free kappa and lambda), 4% had nonsescretory, <1% had IgD. When distribution of stages for our patients according to Durie-Salmon and International staging systems was evaluated, 7% were at stage 1, 23% were at stage 2, 70% were at stage 3. 67% of all cases were at subgroup A, 33% were at subgroup B. According to International Staging System, 27% of patient were at stage 1, 31% were at stage 2 and 42% were at stage 3. Conclusion: This is a retrospective analysis of data from a single center. With data and findings from our study, we hope to gather data from other centers in order to form Turkiye data at near future.
MICROSATELLITE INSTABILITY IN PATIENTS WITH MULTIPLE MYELOMA

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2Süleyman Demirel University, School Of Medicine, Isparta, Turkey

Chromosome 14 abnormalities -mostly translocations- are nearly seen 50 percent of multiple myeloma (MM) patients and these abnormalities are important in the pathogenesis of MM. Genomic instability is a characteristic of tumor cells. Microsatellites are short, tandemly repeated DNA sequences located in genomes. Microsatellite instability (MSI) is the other form to show alterations of DNA mismatch repair system which leads to replication errors. In this report we examined the microsatellite instability in patients with MM in order to point to genomic instability in chromosome 14 and we also analysed 4 different STR locus which are located on different genes. We also compared them with clinical stage and Ig type of disease. We selected 5 different STR loci of chromosome 14 (14q32) and 4 different STR markers named CSF1PO (5q33, 3-34), TH01 (1p15. 5), TPOX (2p25, 1-pter), vWA (12p12-ppter) [Promega Corporation] which are located on different chromosomes. Twenty-six patients were included into the study (10 female, 16 male, mean age 63 year). Seven patients diagnosed as stage one, 7 patients stage two, 12 patients were stage 3. According to Ig type 15 patients had IgG, 5 patients IgA, 5 patients had light chain disease and one had non secretory MM. DNA was extracted from the bone marrow plasma cells after the separation procedure with CD138 magnetic beads (Syndecan-1, Miltenyi Biotec) from the residuæe bone marrow cells and hair DNA was used as control. One of the each pair of PCR primers was fluorescent labeled. Amplified PCR products were run on automatic DNA sequencer (ABI 310, Applied Biosystems) and analyzed using the Genotyper software. MSI was detected in 54% of multiple myeloma patients. Thirty two per cent of patients in D14S65 locus, 25% D14S272, 20% D14S292. Patient samples were also analyzed according to MSI scoring system and 19% had high instability, 35% had low instability 46% of patients had stability at all the loci tested. Sixty per cent of IgG MM had MSI in at least one locus and 27%of all IgG MM patients had high instability, 33% had low instability. We could not find any significant effect of MSI on clinical stage of disease. Five patients with light chain myeloma did not display any abnormality. We could not detect any abnormality on CSF1PO, TH01, TPOX, vWA genes which are located on different chromosomes. As conclusion in present study we showed that MSI is a common finding in MM patients who have heavy chain monoclonal protein especially in chromosome 14q32 region which we know that Ig Heavy chain is being encoded and according to our findings we can also suggest that the molecular defects on this chromosome may lead the malign transformation in MM.

AN UNUSUAL PRESENTATION OF MULTIPLE MYELOMA: PLASMACYTIC ASCITES COMPLICATED BY DUODENAL INVOLVEMENT

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2 Osmangazi University Faculty of Medicine, Department of Pathology, Eskişehir, Turkey  
3 Osmangazi University Faculty of Medicine, Department of Nuclear Medicine, Eskişehir, Turkey

Extradural plasmacytomas are rare, and mostly occur in the upper respiratory tract. Gastrointestinal involvement occurs in only 5% of patients with extradural involvement which includes the stomach most frequently, followed by the jejunum, ileum, colon, rectum, and rarely the duodenum. Peritoneal involvement in multiple myeloma is equally rare. We describe a case of 67-year-old man with Dune-Salmon stage IIIA immunoglobulin A-kappa multiple myeloma, which presented with tense ascites. Abdominal paracentesis revealed atypical plasma cells positive for CD38/CD138 which was confirmed by the presence of a monoclonal peak in the ascitic fluid by protein electrophoresis. The patient’s later course was complicated by gastrointestinal bleeding from a large ulcerated mass localized in the first portion of the duodenum. Biopsy of the duodenal ulcer showed marked monoclonal plasma-cell infiltration by immunohistochremistry. PET study was also performed and high F-18 FDG uptake was noted in the tumor. The patient was successfully treated for bleeding with conservative measures and later underwent VCMPI vincristine, cyclophosphamide, melphalan, prednisolone chemotherapy protocol. Extramedulary spread of multiple myeloma occurs more frequently than is currently recognized. Gastrointestinal involvement may occur soon after the initial diagnosis of multiple myeloma and may be of serious clinical consequence. Failure to recognize myelomatous involvement of gastrointestinal tract may result in achieving inappropriate treatment modalities of surgery, radiotherapy and/or chemotherapy.

TIME INTERVALS PRECEDEING AUTOLOGOUS STEM CELL TRANSPLANTATION (SCT) IN MULTIPLE MYELOMA PATIENTS: A SINGLE CENTER INTENT TO TRANSPLANT ANALYSIS

Mutil Arat, Merih Kızıl Çakar, Ender Soydan, Pervin Topcuoğlu, Aynur Üğur Bilgın, Sule Mine Bakanay, Erol Ayyıldız, Onder Arslan, Muhit Ozcan, Gunhan Gurman, Meral Bekşac, Osman İlhan  
Ankara University, School of Medicine, Department of Hematology, Ankara, Turkey

Introduction&Aim: Hematopoietic SCT activity is less than expected in our country according to 2004 European Activity Survey (1-50/10 million populations). In developing countries many referral centers are facing the problem of having extended waiting lists for transplant candidates. As a referral center with a performance of more than 100 transplants/year we aimed to analyze our kinetics within newly diagnosed myeloma patients under the age of 67, who are de novo transplant candidates. The transplant intervals are calculated in three periods; time from the diagnosis to the PBSC mobilization (tDxMob), time from mobilization to high dose therapy (HDT) (tMobTx) and time from diagnosis to HDT (tDxTx).
Patients: Fifty four multiple myeloma patients under age of 67 admitted to our center between Jan 2004 to Apr 2006 were included into study in intent to transplant approach. The median age was 55 years (32-66 ys) with M/F ratio of 45/15. Results: Autologous SCT candidates (n=43) were followed up for median 18.4 months (1.5-31.7 ms). Forty nine patients were treated with VAD regimen and 5 patients received thalidomide plus dexametasone as first line therapy. After the first line treatment, mobilization was performed in 46 patients (85%) and mobilization (>2x106/kg CD34+ cells/patient weight) was successful in 43 patients (79.6%). The median tDx-Mob was 7.6 months (2.8-27.3 ms). Forty nine patients (74%) received HDT supported by auto-PBSC rescue. We calculated intervals for tDxTx and tMobTx as median 11.7 months (4.6-29.5 ms) and median 2.9 months (0.6-10.2 ms), respectively. Two patients were still waiting for HDT. Unfortunately we have lost a patient, who had been successfully mobilized but died following pulmonary infection before HDT. Eight patients did not enter the mobilization program due to these reasons: Four of them died during primary treatment, two were lost to follow up, another patient is still on treatment and one patient is taken out of the transplant program. Conclusion: In this pilot single centre intent to transplant analysis, we were able to successfully mobilize 85% of the patients. Seventy-nine percent of the patients and 93% of successfully mobilized patients have received HDT. The effects of the deviations of these intervals on the disease-free and overall survival are in further evaluation. This analysis will help us on quality control processes and patient management.

Ref. No: 85 Abstract No: 76

TREATMENT OF RELAPSED/REFRACTORY MULTIPLE MYELOMA WITH THALIDOMIDE: A RETROSPECTIVE EVALUATION IN A CENTER
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1Izmir Atatürk Research Training Hospital, Department of Hematology, Izmir, Turkey 2Izmir Atatürk Research Training Hospital, Department of Internal Medicine, Izmir, Turkey

Between August 2004 and April 2007, 23 cases of multiple myeloma who were refractory to first line chemotherapy or had relapse of disease, and started to thalidomide treatment were evaluated retrospectively. There were 12 men (52.2%) and 11 women (47.8%), total 23 patients, median age was 65 (range 50-77). Eight patients had (34.7%) IgG kappa, 2 patients (8.6%) IgG lambda, 3 patients (13%) undetermined heavy chain kappa, 2 patients (8.6%) IgA lambda, 7 patients (30.8%) IgA kappa and 1 patient (4.3%) was kappa light chain gammopathy. There were 16 patients (69.5%) at stage IIIa, 7 patients (26%) at stage IIa and 1 patient (4.3%) at stage Ia. Median follow up duration was 9 months (3-32 months), median duration of thalidomide therapy was 7 months (2-32 months). Prior to thalidomide therapy 13 patients (56.5%) were treated with VAD chemotherapy, 8 patients (34.7%) with melphalan+ prednisolone therapy, 2 patients (8.6%) with dexamethasone therapy. Also radiotherapy were applied to 8 patients (34.7%) before chemotherapy and 5 patients (21.7%) also treated with oral cyclophosphamide therapy in addition to thalidomide therapy. Median thalidomide therapy dosage were 200 mg (100-300). Side effects of thalidomide were observed in 7 (30.7%) patients; 2 (8.6%) peripheral edema, 2 (8.6%) gastrointestinal side effects, 2 (8.6%) neurologic side effects, 1 (2.3%) pulmonary side effects and 1 (2.3%) tinnitus. In 1 patient (2.3%) thalidomide therapy was stopped due to venous thrombotic event. The result of thalidomide therapy, 1 (4.3%) complete remission, 3 (13.4%) partial remission, 4 (16.8%) minimal response, 11 (47.8%) stable phase disease, 3 (13.4%) refractory to treatment. Also 1 (4.3%) patient was not evaluated due to short term therapy (<2 months). Total response rate was %35.5. In follow up period 4 patients (%17) were died and 4 (%17) patients were out of control. Although two patients who died were not refractory to thalidomide therapy, they had associated cardiovascular diseases that caused death. The estimated one-year OAS for all 23 patients was 65.6%. Our study showed that thalidomide demonstrated appropriate efficacy with acceptable toxicity profile. Influence on patients survival in multiple myeloma patients warrants further studies.

Ref. No: 103 Abstract No: 77

MULTIPLE MYELOMA: RETROSPECTIVE ANALYSIS OF 35 PATIENTS
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Multiple myeloma is a neoplastic monoclonal proliferation of bone marrow plasma cells, characterized by lytic bone lesions, plasma cell accumulation in the bone marrow and the presence of monoclonal protein in the serum and urine. Multiple myeloma accounts for about one
percent of all malignancies. The median age at diagnosis is 65 years and 15% of cases under age 60 years. In this study we evaluated a total of 35 patients with multiple myeloma which was diagnosed according to Kyle-Greipp criteria and made a retrospective analysis regarding clinical characteristics at presentation, outcomes and survival. These patients were followed in our centre from 1999 to 2006. The mean age 64.5 years (range 44-84). Twenty patients (57%) were female and fifteen patients (43%) were male. The diagnosis of all patients had lomber pain, (70%) had fatigue and (12%) had disseminated bone pain in their whole body at presentation. Radiographic skeletal survey was made for all patients. There was no lytic lesion in four patients only but the rest of the patients had multiple lytic lesions. All patients were staged according to Durie – Salmon staging system. Two patients stage I, five patients stage II and two patients were stage III. Twelve of the other twenty six patients were stage IIIA and fourteen patients were stage IIIB. Laboratory findings were as follows: The mean erythrocyte sedimentation rate was 114.5 mm/h (range 62-156) and hemoglobin level 8.7 g/dl (range 11.2-12.3). The mean LDH level and albumin level was 401 U/L (176-1810) and 3.2 g/dl (1.9-4.8) respectively. On the other hand the mean BUN level 89.2 mg/dl and creatinine level was 2.4 mg/dl at presentation. Immunfixation electrophoresis in serum and urine was made for all patients. Serum and urine immunoelectrophoresis confirmed IgG kappa monoclonal gammopathy in five patients, IgG kappa monoclonal gammopathy and kappa free light chain in fourteen patients IgA lambda monoclonal gammopathy in two patients, IgA kappa monoclonal gammopathy in six patients. We have confirmed also kappa light chain in two and lambda light chain in six patients respectively. Twenty six patients received VAD (Vincristine, Adriamycin and Dexamethasone) and nine patients received MP (Melphelan and Prednisolon) for first line therapy. Two patients received VAD after MP. Four patients treated with high dose melphalane chemotherapy and autologous transplant after plateau phase. Refractory or relaps four patients received thalidomide and three patients received bortezomib after VAD. Seven patients received paliative radiotherapy additionally. The mean follow -up duration of patients were 25.4 months. Five patients were out of follow up. Twenty patients were died. The other ten patients are still being follow up. Three years survival rates of stage II 50 % and 27 % for stage III. Since there were no enough cases we have not evaluated patients in stage I.

**Ref. No: 132 Abstract No: 79**

**CLINICAL AND BIOCHEMICAL FEATURES FOR MONITORING MULTIPLE MYELOMA: A RETROSPECTIVE ANALYSIS FROM “DENIZLI LEUKEMIA-LYMPHOMA-MYELOMA STUDY GROUP” (DLLMSG)**

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**Background and Aim:** Multiple myeloma, a neoplasm of plasma cells, accounts for approximately 15% of lymphobematopoietic cancers (LHC) and 2% of all cancers. Incidence rates increase with age, particularly after age 40, and are higher in men than women. “Denizli Leukemia-Lymphoma-Myeloma Study Group” (DLLMSG) was nearly established to register the data of lymphoma, leukemia, and myeloma patients in our city in Western Anatolia. So, we have carried out a retrospective analysis of the clinical and biochemical features of the newly diagnosed multiple myeloma patients followed at our hematology centers. Patients and Methods: Records of all patients in whom multiple myeloma was initially diagnosed at the Departments of Hematology at the Pamukkale University and Denizli Education&Research Hospital from January 2004 to April 2007, were reviewed. Results: Of the 38 study patients, 2. 6% were younger than 40 years, and 25. 6% were 70 years or older. The median age was 65 years (range; 37-78 years). Among patients with multiple myelom 20 (52 %) were male and 18 (48 %) were female. ECOG performance status were ≥2 in 22/38 (58%).

**Ref. No: 132 Abstract No: 79**

**ORAL MELPHALAN AND PREDNISONE PLUS THALIDOMIDE COMPARED WITH HIGH-DOSE THERAPY FOLLOWED BY AUTOLOGOUS PERIPHERAL-BLOOD STEM-CELL TRANSPLANTATION IN PATIENTS WITH MULTIPLE MYELOMA**

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**Background:** High dose chemotherapy followed by autologous stem cell transplant is currently used for the treatment of patients with advanced multiple myeloma. Oral melphalan and prednisone chemotherapy plus thalidomide (MPT) is also an effective treatment in patients who not eligible autologous stem cell transplantation. However, there are no comparative reports of the results of these treatment modalities. Methods: Efficacy of the high dose chemotherapy followed by autologous stem cell transplantation and combination of melphalan, prednisone, and thalidomide have been appreciated in 8 (median age 57; Group I) and 6 (median age 61; Group II) newly diagnosed patients with multiple myeloma, respectively. Results: According to European Bone Marrow Transplantation/ International Bone Marrow Transplantation Registry (EBMT/IBMTR) criteria, 25% of patients achieved immunofixation-negative complete disease remission (CR), 13% achieved a very good partial response, and 62% achieved a partial response, with a 50-89% reduction in monoclonal paraprotein,in group I. Thirtythree percent of patients achieved immunofixation-negative complete disease remission (CR), no patient achieved a very good partial response, and 50% achieved a partial response, with a 50-89% reduction in monoclonal paraprotein in group II. Seventeen percent showed progressive disease in patients who received MPT treatment. The median time to maximum response was 3 months. It was roughly same in two groups. The major acute adverse events (National Cancer Institute Common Toxicity Criteria Grade III-IV) included thrombosis 0% and 16%, infections 12% and 8%, constipation 6% and 50%, and hematologic toxicity 50% and %33 and neurologic 12% and %33 toxicities respectively. Conclusions: These preliminary data suggested that MPT induced rapid and durable tumor responses with CR rates similar to those observed after autologous transplantation. MPT treatment may be suitable first line treatment in myeloma patients. MPT merits further investigation in randomized clinical trials.
patients. Twelve patients (32%) were in stage 3 according to Druie-Salmon staging system. Anemia was present initially in 84% of patients, hypercalcemia (calcium level > or = 11 mg/dL) in 12%, and a serum creatinine level of 2 mg/dL or more in 15%. The beta-2-microglobulin level was increased in 75%. Serum protein electrophoresis revealed a localized band in 28 (73%) of all patients, and immunoelectrophoresis or immunofixation showed a monoclonal protein in 84%. A monoclonal light chain was found in the urine in 66%. Nonsecretory myeloma was recognized in 5% of patients, whereas light-chain myeloma was present in 24%. Conventional radiographs showed lytic lesions in 69%. Extramedullary plasmacytoma were found in four (11%) patients. VAD (Vincristine, Adriablastina, Dexamethasone) were used in 26 patients, and the remaining 12 patients were treated with MP (Melphelane and prednisolone) at initial therapy. Overall response rates were 71%. Ten patients required salvage therapies. Two (5%) of all patients died within 60 days of diagnosis of MM. Infection with renal failure and bleeding are the direct causes of early mortality. Multivariate analysis revealed that age, myeloma cell rate in bone marrow, low platelet count, erythrocyte sedimentation rate, serum lactate dehydrogenase level, serum albumin value, and creatinine value were the most important prognostic factors. The median duration of disease free survival and overall survival were 10 months and 32 months, respectively.

Ref: No: 98 Abstract No: 80

ENDOTHELIAL CELL KINETICS IN PLASMA CELL LEUKEMIA

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Plasma cell leukemia (PCL) is a rare lymphoproliferative disorder characterized by a malignant proliferation of plasma cells in the bone marrow and peripheral blood. PCL is also characterized by a fulminant course and poor prognosis. Despite chemotherapy and the use of novel therapeutic agents patients had short survival. Diagnosis of PCL is established based on Kyle’s criteria which include an absolute plasma cell number comprising greater than 20% of peripheral blood cells. We described one case (53 year, male) of PCL patient. In this case, the bone marrow aspirate smears and biopsy specimens demonstrated a diffuse infiltrate of atypical plasma cells. Immunophenotypic studies showed that case was positive for plasma cell-associated antigens (cytoplasmic immunoglobin, CD38, or CD138) and negative for CD20. We aim to enumerate circulating endothelial cells (CECs) and endothelial progenitor cells (EPCs) during the course of therapy in a patient with primary PCL. A panel of monoclonal antibodies, anti CD146 FITC, anti CD 144 PE, anti CD34 ECD, anti CD117 PC5 were used to enumerate CECs and EPCs CMV pp65 positive patients. Flow cytometric measurement were performed with a FACS calibur flow cytometry (Coulter Epics XL MLC, Beckman Coulter, Florida, USA) equipped with a 15 mW air-cooled 488-nm argon ion laser. Data were analyzed by using EXPO 32 ADC software. CECs number was 33500/mL and EPCs number was 23000/mL before therapy in our patient. After Bortezomib therapy CECs and EPCs numbers were decrease (CECs: 2400/mL and EPCs: 1840/mL). Changes in the number of circulating endothelial cells are becoming prognostic criteria for various clinical events. The quantification of CECs could indicated the presence of endothelial injury, and is simple method of evaluating endothelial-related physiologic and pathophysiologic states. If a correlation between the quantity of CECs and pathologic conditions could be established, circulating CECs could be useful in the diagnosis of the vascular disease, in the explanation of pathophysiologic factors, in the prognostic evaluation of the disease progresses and/or in the evaluation of the treatment efficicany. In one study serum level of VEGF in advanced state of multiple myeloma was elevated and correlated with clinical state. An elevated serum level of VEGF is tought to be associated with a poor prognosis. Plasma cell leukemia represents the most aggressive form of monoclonal gamopathy for which new treatment approaches are needed. Here we report the effect of Bortezomib on both plasma and endothelial cells from one patients with PCL. Bortezomib reduced both plasma cells and CECs numbers. Despite new therapeutic agents, PLC have poor prognosis and short survival. Our patient died 18 month later after diagnosis and we lost him by sepsis and chronic renal failure. These observations may help to improve new therapeutic tool for PCL.

Ref: No:143 Abstract No: 81

OSTEONECROSIS OF THE JAW IN PATIENTS WITH MULTIPLE MYELOMA TREATED WITH ZOLEDRONIC ACID

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Intravenous bisphosphonates; the potent inhibitors of osteoclast mediated bone resorption are one of the most commonly prescribed drugs in the management of multiple myeloma (MM). Zoledronic acid (ZA) is a new generation potent intravenous bisphosphonate, which has been approved for the treatment and prevention of bone lesions, and/or hypercalcemia associated with MM. Osteonecrosis of the jaw (ONJ) is an emerging serious side effect of the new generation bisphosphonates with a growing number of reports related to this pathologic entity. ONJ usually appears following oral surgical and dental procedures, while sometimes spontaneously. These cases are mostly seen and treated by dentists and oral surgeons. The aim of this study was to assess the frequency, characteristics, risk factors and management of ZA induced ONJ in a homogenous group of patients with MM. Twenty six patients with MM who received ZA for a median period of 27 months (min: 5 months, max: 76 months) were evaluated. ONJ was detected in 4 patients and mean drug duration time was 35 months. The frequency was 15.4% and the patients were usually symptomatic. There was no significant difference in terms of the duration of ZA in patients with and without ONJ.

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MYELODYSPLASTIC SYNDROMES

Data from the Registry of the Patients with Myelodysplastic Syndrome from Clinic of Hematology, Fundeni Clinical Institute, Bucharest, Romania. I. Epidemiological General Data

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Background. Since the World Health Organization (WHO) recognized MDS as a disease entity only starting with 1997, epidemiological data on MDS cannot be obtained from official statistics on morbidity and mortality and have to be extracted from specialized registers. We present the first Romanian study on the incidence and characteristics of MDS, based on the data existing in Fundeni Clinical Institute, Bucharest, the greatest hematological department in Romania. Method. The MDS files at diagnosis of the patients admitted during the period 1980-2005, recorded in the registration forms provided by the MDS Foundation (USA), represented the primary data-base. The hematological data of the MDS patients included in the registry were re-evaluated and classified according to French-American-British (FAB) criteria. The distribution by sex, age groups, subtypes and the annual number of new cases were analysed comparatively with other reference studies. Results. Four-hundred and three cases of MDS were identified. The distribution between sexes was relatively balanced with a slight global preponderance of males (M/F 1.26), except for refractory anemia with excess of blasts (RAEB) 1. 94. The mean age at diagnosis was 62. 3 years (16-90). Most of the patients (60. 6%) belonged to the group of age 61-80, where all the subtypes of MDS had the highest rates. A noticeable proportion (17%) had ages below 50 years, 25% of which in the range 16-30. On the other hand, few cases (4%) were above 81. Patients with refractory anemia (RA) and refractory anemia with ringed sideroblasts (RARS) accounted for 44. 5% of all cases (RA 29%, RARS 15. 5%), RAEB and RAEB in transformation 33%, chronic myelomonocytic leukemia 5. 6% and unclassified 16. 7%. The annual number of new cases was constantly low during the period 1980-1989, but increased dramatically from 11 cases/year in 1990 to a maximum of 48 cases/year in 1999, showing a clear decrease afterwards. The subtypes with the most important increase in time were RA and RARS. Conclusions. This study indicates an actual increase of the number of MDS cases in Romania over the investigated period of time. Particularly, a noticeable proportion of young patients and a low proportion of patients ≥ 81 years have been found, which make our findings in the middle between the Asian than to the Western MDS epidemiological results. e-mail: mds.fundeni@yahoo.com

Treatment of High-Risk Myelodysplastic Syndrome with Demethylating Agents

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Myelodysplastic Syndrome (MDS) comprises a heterogeneous group of clonal hematopathies derived from an abnormality affecting a multipotent hematopoietic stem cell and characterized by maturation defects resulting in ineffective hematopoiesis. It most frequently occurs in elderly patients. Despite trials testing numerous agents in patients with MDS, no single drug has yet emerged as an accepted standard of treatment. The effect of available lineage-specific growth factors is limited to improvement of single lineages and has not resulted in the survival benefit. Observation and supportive care with blood products and antibiotics, when necessary, continue to be the mainstays of therapy. We administered 5-azacytidine, a cell-cycle specific ring analog of the pyrimidine nucleoside cytosine, as a continuous intravenous infusion, 75 mg/m2 per day for 7 days every 4 weeks to two MDS patients, one of whom is 48-year-old-female and the other is 77-year-old-male. The patients had refractory anemia with excess blasts (RAEB) and refractory anemia with excess blasts in transformation (RAEB-T). One patient was received two cycle of 5-AZA and the other was received only one cycle. During the observation period after the treatment, a clear hematologic response, decrease in the need of transfusion and blasts clerereance did not occur. Hematological toxicity was mild and consisted of thrombocytopenia and leukopenia. Extramedullary toxicity consisted of arthralgia, diarrhea. But both of the patients died in thirty days due to sepsis. As the result of our observation we may suggest that after this treatment the control of the disease is hard and the possibility of infection is frequent.

Flow Cytometric Analysis of Peripheral Blood in Diagnosis of Myelodysplastic Syndromes

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Myelodysplastic syndromes (MDS) are a heterogenous group of diseases and more objective tests are needed for diagnosis. Flow cytometric analysis of peripheral blood has been suggested useful for diagnosis of MDS, recently. In this study we evaluated the diagnostic value of peripheral blood analysis by flow cytometry in MDS. 16 patients with MDS, 5 patients with cytopenias other than MDS and 10 healthy controls are included in the study. Peripheral blood samples are stained with CD45, CD33, CD7, CD13, HLA DR, CD117, CD34, CD10, CD11b, CD11c, CD71, CD16, CD15, TdT, MPO monoclonal antibodies and analysed at flow cytometry (BD-FACS Calibur). Neutrophils, lymphocytes and monocytes are
identified by CD45/SSC. Presence of abnormal cell population, positive or negative antigenic expression or different expressions, hypogranularity are evaluated. Abnormal cell population (7/16), hypogranularity (1/16), CD10 negativity (8/16), presence of HLA DR (3/16), presence of CD117 (3/16), lack of CD33 expression (2/16), lack of CD13 expression (2/16), presence of CD34 (2/16), lack of CD11c (2/16) on neutrophil gate; presence of CD33 (10/16), presence of CD13 (8/16), presence of CD117 (3/16), presence of CD34 (1/16) on lymphocyte gate were most frequent abnormalities in patients with MDS. In conclusion, flow cytometric analysis of peripheral blood is useful in MDS diagnosis. Most common abnormalities are presence of abnormal cell population on CD45/SSC, lack of CD 10 on neutrophils, presence of CD33 and CD13 in lymphocyte gate.

**MYELOPROLIFERATIVE DISORDERS**

Ref: No: 97 Abstract No: 85

**CIRCULATING CD34 CELLS IN MYELOFIBROSIS**

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Myelofibrosis with myeloid metaplasia was first described in 1879, classified as a myeloproliferative disorder in 1951, and characterized as a clonal stem cell disorder in 1978. It characterized by pancytopenia with intact maturation, progressive bone marrow fibrosis, and multiorgan extramedullary hematopoiesis. Idiopathic myelofibrosis characterizes may lead to an increased number of circulating CD34 positive cells in the peripheral blood. We aim to absolute number of circulating CD34 positive cells by flow cytometry in patients with idiopathic myelofibrosis. The diagnosis criteria of myelofibrosis were utilized for the Italian Consensus Conference criteria. We enrolled 11 patients (6 women and 5 men; age range, 45-74 years) with myelofibrosis. The patient group was a non-selected group. Peripheral blood samples were drawn into EDTA anticoagulated tubes. A panel of monoclonal antibodies, anti CD10 on neutrophils, presence of CD33 and CD13 in lymphocyte gate. An analysis of pooled outcome data from 461 pregnancies in ET showed that the average platelet count was 1010x10^9/L in patients with successful pregnancies. Our patient had normal platelet levels during her pregnancy. The mechanism responsible for such decline is unclear. This observation suggested that hematological effects of pregnancy may show individual discrepancies.
100%. Four of the patients stated that they drank BR solution roughly 200 mL/week. However, we found that our participants with CML had no severe symptoms or other clinical manifestations of fluid retention at the time of the study. This observation did not support the thesis that BR solution might induce fluid retention in patients with CML using imatinib.

Monitoring response at molecular level is suggested to provide a new reference measurement in treatment of Philadelphia chromosome positive chronic myeloid leukemia (Ph+ CML) and plays an important role in optimizing treatment. The molecular response is described as removal or reducing of Bcr-Abl transcripts which produce abnormal proteins responsible for white blood cell proliferation in CML patients. In these patients who showed a major molecular response along with a complete cytogenetic response at 12th month, the survival ratios, without progression, were expressed as 100% at 24th month. With studies showing that Glivec (imatinib) provides a better molecular response than conventional combination therapies, the molecular response is expressed to be a new reference measurement that can reveal success of CML treatment. However, long term results of imatinibe in CML patients are not well known. In this report, we presented findings of 6 Ph+ CML cases (3 female, 3 male, all younger than 50 years of age) having received imatinib therapy between the years 2005 and 2007. For four of the patients, exactly matching donors were found but for one of the patients the search of a donor had not been completed yet. According to Sokal scoring 2 patients were located in low-good, 3 patients in intermediate and 1 patient was in high-poor risk group. The hematological response was observed at 12th day in the low risk group and approximately at 45th day in the other groups. On the other hand, cytogenetic response was seen in 3rd month in low risk patients and approximately in 9th month in intermediate and high risk groups. Hematological and molecular responses were achieved in all patients in one year. In 24th month, Philadelphia chromosome became positive in PCR in one patient from poor risk group. Generally imatinib therapy was well tolerated. However, since we do not know the long term outcomes, there is not sufficient data about the question “Which patients should receive PSCT first?”

Today, we know that the curative therapy can only be achieved with PSCT. Particularly in patients from early chronic phase and good risk group, long term disease free and total survival was observed with PSCT. However, factors affecting transplantation related deaths must be described well because of the long term mortality and morbidity that are associated with stem cell transplantation. But we still think that PSCT will be a good treatment alternative in cases with HLA matched sibling donor and at early chronic phase with molecular and hematological response, particularly in young-poor risk patients (<45 years).

PERIPHERAL POLYNEUROPATHY ASSOCIATED WITH IMATINIB TREATMENT
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It has been reported that neurological side effects and complications peripheral may occur during the course of chronic myeloproliferative diseases. Among these complications, peripheral polyneuropathy has been shown to be associated with the development of cryoglobulinemia, paraproteinemia and/or interferon therapy. Imatinib (a tyrosine kinase inhibitor) was FDA approved because of its exceptional safe profile. However, polyneuropathy associated with imatinib has not been reported yet. A 58-year-old man was hospitalized because of leukocytosis. A diagnosis of chronic phase chronic myelocytic leukemia was made. The patient did not complained of dysesthesia or pain in his feet during the initial therapy of hydroxyurea lasting after four weeks. Administration of imatinib were started. Two months later, the patients achieved hematologic response. After 8 weeks of imatinib treatment, the patient complained of worsening dysesthesia an severe pain in his feet, particularly in the right. An mixed form of peripheral neuropathy was diagnosed by electrophysiological examination. The patient had no history of diabetes mellitus, hypertension, hyperlipidemia, and alcohol consumption. Biochemical and immunological studies revealed no paraprotein and cryoglobulin. There was no sign of vasculitis. His neuropathic complaints persisted for two years Therefore, the clinical course suggested that mixed cryoglobulinemia was associated with chronic myelocytic leukemia.

FEVERILE NEUTROPENIC EPISODES IN ACUTE LEUKEMIA PATIENTS: EXPERIENCE OF BAŞKENT UNIVERSITY HOSPITAL
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Feverile neutropenia is a major cause of morbidity and mortality in acute leukemia patients. Besides infections caused by resistant bacterial microorganisms, fungal and viral infections are serious clinical problems in these patients. Although there are guidelines for management of the febrile neutropenia, every hospital has its own microorganism spectrum and every center has to know its own patient profile. Social securities, costs, clinical experiences and patient’s own clinical characteristics effect the antimicrobial choice. With this study we decide to reveal our data so that we can make more accurate decisions in future febrile neutropenia episodes and can compare our results with other centers and we can also let them to do so. We researched the medical files of our adult acute leukemias admitted to our hospital since January 2002. 42 febrile neutropenia episodes of 22 patients were taken under consideration. Mean age was 48 (SD±19) years. There were four acute lymphoblastic leukemias (ALL), two secondary leukemias, 16 acute myeloid leukemias (AML). Fever developed minimum at day 0 of absolute
neutropenia and maximum at day 15 (mean 3. 19 days). Most of the febrile episodes were on day 0 (16 of 42) and most of these first day fevers were newly diagnosed patients (68. 8%). At diagnosis 69% of patients were started treatment without any signs of infection other than fever, 14. 3% of patients had clinical sign of infection, at 11. 9% of patients radiology effected the treatment, only 2 episodes (4. 8%) there had been culture positivity guiding the initial therapy. On follow up, radiologic tests showed infection’s source in 31% of episodes and at 38. 1% of the episodes, the agent was isolated. 11 episodes (24%), had proven fungal infection. At 15 (47%) of 32 febrile episode with no proven fungal infection, some kind of antifungal agent were used and in 11 (73%) of them liposomal amphoterin B was the choice. On 27 episode (64. 3%) GCSF was used. In patients receiving standart or high dose treatment, GCSF was used in 67. 7% of the episodes, with reduced dose regimens the percentage was 71%. Among all febrile episodes 71. 4% ended with resolution, 14. 3% ended with death, 6 patient (14. 3% of episodes) were discharged because of the resistancy of their primary illness. In our study in 31% of patients we couldn’t show the infectious source of fever in the follow up. Empirical antifungal therapy for suspected infections is standart of care in neutropenic cancer patients with persistent fever despite broad-spectrum antibiotics. In our center in about half of the episodes, antifungal treatment is added to the therapy. GCSF use in our patient group is parallel with the literature. Sample of cases are insufficient to give complicated statistics but it revealed our patients data and will be used in our future desicions. On this very important subject we need more data and future studies will be planned.

### STEM CELL TRANSPLANTATION

**Ref No: 7**

**Abstract No: 91**

**TESTS, HISTORICAL EFFORTS AND IMMUNE RECONSTITUTION IN CORD BLOOD STEM CELLS TRANSPLANTATION**

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Allogeneic stem cell transplantation is an accepted treatment modality for selected malignant and non-malignant diseases. However, the ability to identify suitably matched related or unrelated donors can be difficult in some patients. Alternative sources of stem cells such as cord blood provide a readily available graft for such patients. Since the cell numbers of hematopoietic progenitors in cord blood is limited and the collection can occur only in a single occasion, its use in adult patients can be more problematic. The patient outcomes should be review and analyze for various factors such as cell dose, HLA typing, and patient selection that could have contribute to the final outcome of these adult patients. Discussion of the various benefits and risks should be present. Description of the historical efforts associated with expansion of hematopoietic stem cells, specifically with cord blood cells expand cord blood cells continue with novel methods. Moreover, a better understanding of stem cell biology and signaling is critical if we are to be able to effectively expand these cells for clinical use. Describe the immune reconstitution or lack thereof following cord blood transplantation appears be very important, one of the hallmarks of successful hematopoietic stem cell transplantation is the ability to fully reconstitute the immune system of the recipient. Thus, the relationship between stem cell source and the development of T lymphocyte functions required for protection of the recipient from infection will be described, and cord blood recipients will be compared with those receiving other sources of stem cells. Key words: stem cell, cord blood, transplantation

**Ref No: 70**

**Abstract No: 92**

**AUTOLOGOUS HEMATOPOIETIC STEM CELL TRANSPLANTATION FOR LYMPHOMA AND MYELOMA AT YEDITEPE UNIVERSITY HOSPITAL**

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Stem cell transplantation(SCT) is an effective treatment modality in hematological malignancies. SCT Unit at Yeditepe University Hospital(CIC-919) was activated in October 2005 and accredited for unrelated SCT by EBMT in April 2007. Until April 2007, a total of 26 autologous transplantation was performed for 18 patients with lymphoma (9 NHL, 9 HL) and 8 patients with myeloma. Patients with lymphoma received high-dose sequential chemotherapy consisted of HDVP16 followed by HD-MTZ+HD-MEL preparative regimen after IIVP or HIDAC+HDXTX salvage regimens. Patients with myeloma received HD-MEL at 200 mg/m2 if they are less than 70 years of age, and 140 mg/m2 over 70 years of age. Median age was 43,5 (13-71) years and mean time from diagnosis to transplant was 2,1 years. Lymphoma patients received a mean number of 8 (2-18) salvage regimens prior to transplantation. All patients engrafted and median engraftment period was 13 (9-24) days. Median follow-up period was 7 (1-18) months. During follow-up of 18 patients with lymphoma, 4 patients (22%) relapsed and one died during the follow-up period. 2 of these patients are induced into CR, one following allogeneic transplantation and the other following radiotherapy combined with R-EPOCH regimen. DFS for lymphoma patients at 18 months is 73. 3% at 18 months. Of 8 patients with myeloma, 2 patients (%25) relapsed, and one patient was found to be refractory to HD-MEL preparative regimen. The patient with refractory disease is currently in CR following an unrelated allogeneic stem cell transplantation. Of 2 patients with progressive disease, 1 is currently in CR following 2 cycles of EPOCH regimen and the other died secondary to progressive disease. Post-transplant complications were DVT, transient hyperbilirubinemia, sleep apnea related cardiac arrest which responded to resuscitation (n=1 each). A patient with myeloma experienced transient congestive heart failure in and HBV hepatitis following a dental procedure. All patients were alive following auto-SCT at day +100 (TRM=%0), only one patient with myeloma died due to relapse at 9 months. At present, of 25 surviving patients, 20 are assessed for response and 18 of these patients are in CR. The high survival rate (96%) achieved following autologous SCT, low TRM (0%) and the high remission rate (90%) may be related to team approach, 24-hour patient follow-up, and effective management of complications encountered during early transplant period. Patients with lymphoma or myeloma relapsing following autologous transplantation can be successfully salvaged by a second allogeneic transplantation procedure.
Background: Insulin resistance syndrome has been shown to be associated with many coagulation and fibrinolytic proteins and these associations suggest that some coagulation and fibrinolytic proteins have a role in atherothrombotic disorders. Aim: This study was conducted to determine the levels of some of the haemostatic parameters in subjects having metabolic syndrome and to correlate these values with the anthropometric and metabolic variables associated with this syndrome. Subjects and methods: The study included 46 obese non diabetic subjects of whom 28 subjects (group 1) fulfilled the ATP III criteria of the metabolic syndrome and 18 subjects (group 2) did not have metabolic syndrome as well as 14 lean subjects (group 3) of matched age and sex as a control group. Clinical and laboratory evaluation of the study groups stressed on anthropometric measurements (weight, height, body mass index, waist circumference, and sagittal abdominal diameter), blood pressure, and laboratory measurements of fasting plasma glucose, fasting insulin, serum lipids, tissue plasminogen activator (t-PA), antithrombin III activity (ATIII), protein C and von Willebrand factor (vWF) antigen. Results revealed: Significant increase in the concentrations of t-PA and vWF antigens in subjects having metabolic syndrome (group 1) in comparison to the other groups while there were non-significant changes in the levels of protein C antigen and AT III activity. Both t-PA and vWF showed significant correlation with HOMA-IR as a measure of insulin sensitivity. The t-PA showed also significant correlation with most of the variables of metabolic syndrome including waist circumference, BMI, systolic blood pressure, fasting plasma glucose, fasting insulin, and HDL cholesterol. On the other hand, vWF showed significant correlations with fasting plasma glucose, fasting insulin and sagittal abdominal diameter, with non-significant correlations with the other variables. Conclusion: Haemostatic and fibrinolytic parameters should be included in the features and characterization of the insulin resistance syndrome. t-PA and vWF antigens concentrations were increased in subjects with metabolic syndrome and correlated with the HOMA-IR measure of insulin sensitivity. Taking into consideration that both t-PA and vWF are mainly released from vascular endothelium, these findings could be an indicator of endothelial dysfunction in that group of subjects.