14th Congress of the
European Society for
Haemapheresis and
Haemotherapy

Abstracts

Prague, September 10–13, 2003

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The 14th Congress of the Interdisciplinary European Society for Haemapheresis and Haemotherapy (ESFH) was held on September 10–13, 2003 in Prague, in association with the 9th Working Days of the Czech Society of Transfusion Medicine and the Slovak Society of Haematology and Transfusion Medicine. It was the first time that the Czech Republic hosted an international congress on hemapheresis.

The standard of health care in the Czech Republic has always been high. After the ‘Velvet Revolution’ and the democratization in 1989, a rapid development of hemapheresis and hemotherapy started. Currently, there are more than 250 cell separators operating in the country, which is quite a figure in proportion to population. Donor and therapeutic apheresis procedures have grown in number and types, including not only PBPC harvesting and DLI, but also photopheresis and immunoapheresis.

For the hosting Czech Transfusion Society, the 14th congress of the ESFH was a most welcome international forum to draw the attention on their activities and to demonstrate the quality and importance of their research. It was the aim of the International Scientific Committee to present the different aspects of hemapheresis, reflecting research and technical aspects as well as future developments in this rapidly developing field. A main focus of the conference was the emergence and establishment of novel hemapheresis methods and indications in the 2 years since the previous ESFH congress.

The scientific program consisted of 74 oral presentations (36 invited speakers) as well as 71 posters, with contributions not only from hemapheresis specialists, but also from nephrologists, intensive care specialists, and health professionals from adjacent branches of medicine, reflecting the interdisciplinary character of the conference and the ESFH. The congress included also educational programs and a special session for Czech and Slovak technicians. More than 540 physicians and health care specialists (including 113 from the Czech Republic) participated in the 14th ESFH congress.

During the congress, the Hoegman Award, which is announced every 2 years to honor the best young scientist in the field of hemapheresis, was given to Dr. Pavel Žák, Ph.D., from the Department of Internal Medicine II – Hematology, University Hospital in Hradec Králové. Dr. Žák is an active member of research groups dealing with erythrocytapheresis, plasmapheresis, immunoapheresis (LDL apheresis) and photopheresis, and was honored for his optimization of venous access for hemapheresis.

This volume includes a selection of the summaries of the outstanding educational lessons, oral presentations, SAFs (Seminars of Advances in the Field) and posters. For easier orientation, the summaries are numbered in the same way as the papers, oral presentations or posters in the Abstract Book. However, this selection cannot cover the entire spectrum of apheresis presented during the conference. Some copies of the Abstract Book including all contributions of this year’s congress are still available for those who are interested (ISBN 80-903238-8-X, publisher: HK CREDIT, Ltd., Hradec Králové, Czech Republic).

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Editorial

Hemapheresis in the Czech Republic

Short Comments to the 14th Congress of the Interdisciplinary European Society for Haemapheresis and Haemotherapy, Prague, September 10–13, 2003)
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Non-infectious complications of HSC transplantation
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Spleen Size Changes in Peripheral Blood Hematopoietic Progenitor Cell Donors Given G-CSF

D.F. Stroncek, S.F. Leitman
National Institute of Health, Dept. of Transfusion Medicine, Bethesda, USA

Background: Granulocyte colony-stimulating factor (G-CSF)-mobilized peripheral blood progenitor cells (PBPC) are replacing marrow as a source of hematopoietic progenitors for transplantation. PBPC donors given G-CSF experience splenic enlargement and rarely, spontaneous rupture of the spleen. This study evaluated the incidence and time course of splenic enlargement in PBPC concentrate donors and assessed factors affecting size changes.

Methods: Healthy adults were given G-CSF (10 micrograms/kg/day) for 5 days and a PBSC concentrate was collected by apheresis. Ultrasound was used to assess cranio-caudal spleen length prior to giving G-CSF and the day of apheresis. In one group of donors was measured again 3 or 4 days after apheresis (n = 20) and in a second group of donors spleen size was measured again 10 days after apheresis (n = 5). The effects of donor age, gender, race, and changes in blood chemistries, blood counts, and CD34+ cell counts on spleen length change were assessed.

Results: Among donors in the first group spleen length increased in 19 of 20 donors. Mean length changed from 10.9 ± 2.0 cm pre-G-CSF to 12.3 ± 2.1 cm on the apheresis day (p < 0.001). The mean increase in length was 1.5 ± 0.9 cm (13.3 ± 9.1%). Spleen length increased 20% or more in 6 subjects. Three or four days after apheresis the spleen length fell to 11.3 ± 1.8 cm (p < 0.001), but it remained greater than baseline levels (p = 0.003). Spleen length change was not affected by donor gender, race, or age. There was no relationship between changes in spleen length and baseline and apheresis-day blood counts and chemistries and changes in blood counts and chemistries. In the second group of donors, spleen length increased in all 5, but returned to baseline levels 10 days after apheresis. There was no difference in spleen length measured before G-CSF and 10 days after apheresis (10.3 ± 1.2 cm versus 10.0 ± 1.4 cm).

Conclusions: Spleen size increases in almost all PBSC donors. Enlargement is transient and brief but marked in some donors and may place them at risk for splenic rupture.

O 13 Influenze of Donor Characteristics and G-CSF-Administration Schedule on the Efficacy of Peripheral Blood Progenitor Cell Mobilisation

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1Transfusionsmedizin, Medizinische Klinik und Poliklinik I, Klinikum der Technischen Universität, Dresden, Germany

Purpose: CD 34+ cell mobilisation in healthy donors varies to a wide scale. Defining predictive factors of mobilisation efficacy is of great interest to optimise protocols for allogeneic stem cell donors.

Methods: 1474 healthy donors (986 men, 488 women) underwent G-CSF application and PBPC collection at our department between 1/1996 and 8/2012. G-CSF administration was performed in 3 dosages: filgrastim 10µg/kg/day on 5 days; lenograstim 7.5µg/kg/day on 5 days; lenograstim 7.5µg/kg/day on day 1 and 2, 12.5µg/kg/day on the following 3 days. Leukapheresis was performed at day 5 (and 6, if necessary). CD34+ concentration in peripheral blood (×10^11/L) at day 5 before 1st leukapheresis was analysed for correlation with the following parameters: leukocyte and platelet counts before G-CSF administration, sex, age, body mass index (BMI), nicotine and alcohol consumption of the donors, G-CSF dose and mode of G-CSF application (single dose versus split dose).

Results: The median concentration of CD 34+ cells in peripheral blood at day 5 was 56±10 µl in males and 42±10 µl in female donors (p < 0.0001). A 2nd apheresis had been performed in 34% of males and 53% of female donors. A significantly positive correlation of CD 34+ concentration was found with BMI (p < 0.001) and the schedule of G-CSF application (split versus single dose: p < 0.0001). In a multivariate analysis, the schedule of G-CSF application had the most significant influence on the efficacy of peripheral blood progenitor cell (PBPC) mobilisation.

No significant correlation was found with G-CSF-dose, donor age, alcohol consumption and smoking status.

Conclusions: In our donor population PBSC mobilisation worked best in male donors with higher BMI. The schedule of G-CSF-administration seems to be very important for the mobilisation efficacy in healthy volunteer donors. Dose splitting of G-CSF should be performed, whenever possible.

O 14 Extracorporeal Photochemotherapy (ECP) in the Treatment of Acute and Chronic GvHD: Possible Relationship with the Number of Treated MNC Cells

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Servizio Trasfusionale, Ospedale San Gerardo-Monza e
*Clin. Pediatria, Univ. Milano Bicocca

Background: ECP proved its effectiveness in patients either with aGvHD or cGvHD refractory to standard immunosuppressive treatments. Nevertheless, there are still at least two controversial issues: 1) what is the optimal treatment schedule and 2) does a correlation exist between the number of collected/irradiated MNCs (lympho-mononuclear) and clinical response? We retrospectively analyzed our pts series to elucidate this latter issue.

Patients and Methods: 26 pts who underwent ECP for aGvHD (6 pts) or cGvHD (20 pts) were studied. Clinical data (grading, organ involvement, clinical response, etc) were recorded as well as the number of MNC/Kg collected, irradiated and reinfused in each ECP procedure. MNC collection was performed by using a Cobe Spectra device (standard WBC collection set). 8-MOP was added at 200 ng/mL final concentration, UV-A irradiation (2 J/cm^2) was performed by using the Uvamat device. Data are shown as median and range.

Results: 393 ECPs were performed over a 5-yr period; 71 in aGvHD pts, who underwent 10 (6-14) ECPs each and 321 in cGvHD pts who underwent 18 (2-29) ECPs each. Overall aGvHD grading ranged from grade I to grade III, while 16 out of 20 pts had extensive cGvHD. Skin was the major organ involved in 5 out of 6 aGvHD pts and in 18 out of 20 pts with cGvHD; 8 pts from this latter group also had mucosal involvement. Basal pre-ECP WBC count was 2.5 (1.1-15.1) × 10^9/L in aGvHD pts and 5.6 (1.4-15.6) × 10^9/L in cGvHD pts. Clinical response to ECP was as follows: 4 CR, 1 PR and 1 NR in aGvHD pts; 10 CR, 6 PR, 3 NR and 1 NV (two ECPs only) in cGvHD pts. Individuals with cGvHD who showed CR or PR received a double MNC dose compared to NR: 115 ± 10^6/Kg vs. 64 ± 10^6/Kg. On the contrary no difference was detected in aGvHD pts.

Conclusions: It has been postulated that ECP might trigger a specific APC-mediated response against auto-reactive lymphocytes or induce apoptosis in collected and irradiated MNCs. Our report suggests a possible relationship between MNC dose and clinical response, at least in cGvHD. Further studies on more pts are needed to ascertain the importance of MNC dose in treating GvHD.
finally, UV-A irradiation (2J/cm²) was performed by means of a Uvamatic (Vilber-Lourmat) device.

Results: All pts completed the study. MNC collections were performed without clinically relevant side-effects but mild hypocalcemia-related symptoms. Pts were given 5.6–11.1 x 10⁹ irradiated cells/ECP (median dose). PB WBC count did not significantly change all over the study period, as well as CD4+, CD8+ lymphocyte count and CD4/CD8 ratio. Three pts had no relapses, and two had 3 relapses each, responsive to low dose steroids. In 2 pts steroid total dose tapering was achieved and three did not required steroids all along the study period. EDSS did not worsened in 4 out of 5 pts. MS activity evaluated by means of MRI was reduced compared to the pre-ECP period.

Conclusions: ECP proved to be feasible and well tolerated in all pts; a substantial lowering or withdrawal of steroid therapy was obtained in all cases. We observed a decrease in the number of relapses, whilst EDSS stabilized in 80% of the pts. Similar findings were observed as to the occurrence of new cerebral lesions in MRI examination after gadolinium enhancement. A prospective multicenter study is now required to assess the possible usefulness of ECP in the treatment of RR-MS.

L 21
PBPC Collection Techniques: Is There a Way to Get enough CD 34+ Cells in ‘Poor Mobilizers’?
Z. Gaslová, J. Hrusňková, J. Žlabová, I. Marinov, Š. Vodvárková
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Introduction: PBPC collection techniques and subsequently the yield of CD 34+ cells play an important role in the effect of autologous and allogeneic transplantation. PBPC are usually collected in the standard or in the large volume leukapheresis regime (LVL), but a criterion for selection of the optimum method has not yet been defined. The effect of mobilization and consequently the clinical condition of the donor or patient could affect directly the choice of the collection technique.

Methods: We evaluated the results of 88 standard and 217 LVL collections in 46 healthy donors and in 149 patients who suffered from hematologic oncological diseases. More than 3 total blood volumes of the donor or patient were processed in LVL procedures. The precollection concentration of CD 34+ cells in blood equal or higher than 20 x 10⁶/ml was considered as a criterion of a good effect of mobilizing therapy.

Results:

<table>
<thead>
<tr>
<th>Collections</th>
<th>Yield of CD 34+ cells (10⁶/kg)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Standard</td>
<td></td>
</tr>
<tr>
<td>LVL</td>
<td></td>
</tr>
<tr>
<td>Donors well mobilized</td>
<td>4.1 (1–13.6)</td>
</tr>
<tr>
<td>Patients well mobilized</td>
<td>3.1 (1.9–43.2)</td>
</tr>
<tr>
<td>Patients weakly mobilized</td>
<td>0.7 (0.2–1.2)</td>
</tr>
</tbody>
</table>

LVL enabled to get a higher yield of CD 34 cells than the standard collections in well mobilized donors, in well mobilized patients as well as in weakly mobilized patients. The yield of CD 34+ cells equal or higher than 6 x 10⁶/kg was obtained from a single LVL procedure (Cobe Spectra) in donors and in well mobilized patients.

Conclusion: We can recommend LVL in all donors or in patients who can tolerate it. LVL should also be preferred in weakly mobilized patients where the chance to collect at least a minimum amount of CD 34+ cells for transplantation is possible using only this technique. No serious adverse reactions in LVL have been observed so far.

O 27
Efficacy of Cytoreductive Leukopheresis in Acute Leukemia Patients with Hyperleukocytosis
Ankara University Faculty of Medicine, Dept. of Hemapheresis Unit and Blood Bank, Ankara, Turkey

Objectives: Therapeutic leukopheresis is used to control the acute symptoms of hyperleukocytosis by cytodestruction. The symptoms of hyperleukocytosis are usually prominent when the leucocyte count is above 75 x 10⁹/L among acute myeloblastic leukemia (AML) patients. Because mortality increases significantly when WBC counts reach higher levels than 100 x 10⁹/L, there is an indication for therapeutic apheresis to prevent tumor lysis and hyperleukocytosis syndrome. Herein we report the procedure based results of 30 acute leukemia and chronic myeloid leukemia (CML) in blastic phase (BP) patients who underwent cytoreductive leukopheresis between January 2000 and September 2002.

Methods: The median age was 41 (16–74) years, male to female ratio was 23/7. The diagnosis of the patients were AML 20 (66.7%), acute lymphoblastic leukemia: 5 (16.6%) and CML in BP (16.8%). The venous access was supplied by peripheral vein in 80% (n = 24) of patients and by central venous catheter in the remainder (20%, n = 6). Baxter Fenwall CS3000+ (n = 30) and Cobe Spectra devices (n = 1) were used. Leukopheresis was performed in median 1 (1–4) session and median 7L (4.5–9.5) L blood was processed in median 170 (135–220) minutes. Erythrocyte and platelet transfusion were performed if necessary after the procedure.

Results: The median leucocyte count, platelet count and hemoglobin levels of patients before and after leukopheresis were 176 x 10⁹/L (93.8–459) – 83.4 x 10⁹/L (58–733), 60 x 10⁹/L (10–750) – 45 x 10⁹/L (11–457) and 8.5 g/dl (6.7–15.7) – 8 g/dl (5–14.2), respectively. The median decrease in the white blood cell count was 45.8% (1.8–81.6). One patient (3.8%) failed to achieve an effective cytoreduction. There was only one hypocalcemic case who required replacement (n = 1), procedure was ceased because of a life threatening dyspnea and angina in a patient during the third cycle (n = 1), no patient died during the procedure. Leukopheresis is an effective method to achieve a rapid cytoreduction. The administration of hydroxyurea (4–6 g/day) po accompanying the procedure supposed to be more effective than the procedure alone.

Conclusions: In spite of low preapheresis platelet counts, successful apheresis could be performed without major complications. There was no evidence in the efficacy of therapeutic leukopheresis between AML, ALL, CML groups. The effects of this procedure in the remission rates and the survival should be investigated and compared retrospectively with the cases whom leukopheresis was not performed.

<table>
<thead>
<tr>
<th>AML</th>
<th>ALL</th>
<th>CML</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number of patients (n)</td>
<td>20</td>
<td>5</td>
</tr>
<tr>
<td>Median age</td>
<td>34</td>
<td>50</td>
</tr>
<tr>
<td>Median number of procedure</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>Median procedure time (min)</td>
<td>180</td>
<td>160</td>
</tr>
<tr>
<td>Median decrease in WBC (%)</td>
<td>44.28</td>
<td>51.8</td>
</tr>
</tbody>
</table>

L 37
Treatment of Thrombotic Thrombocytopenic Purpura with Plasma Exchange Using Fresh Frozen Plasma or Cryosupernatant Plasma: Variations in Metalloprotease and vWF Levels
G. Rock and the Canadian Apheresis Group
The Ottawa Hospital, Hematology, Ottawa, Canada

Introduction: TTP remains a catastrophic thrombotic disorder which, because of various forms of plasma therapy, now results in recovery in up to 90% of patients. Current therapy is based mainly on plasma exchange, using either FFP or CSP. Data from several trials have indicated that CSP, which is deficient in the higher molecular weight multimers of vWF, is better therapy than FFP. The pathophysiology of the disease has long been an enigma: in 1998 both Tsai and Furlan reported a deficiency of...
a WVF cleaving metalloprotease in congenital and some acquired forms of TTP.

Methods: We have determined outcomes and metalloprotease levels in 50 patients randomly assigned to receive 1.5 volume plasma volume exchange with either FFP or CSP.

Results: 25/26 patients receiving CSP and 22/24 of those on FFP were alive at 1 month. Prior to treatment, VWF levels ranged from 1.0 to 3.8 IU/ml. After treatment this decreased significantly with the level considerably lower in patients receiving CSP. The metalloprotease levels showed large variation ranging from 10–100% protease activity. None of the patients demonstrated protease activities of <5% of control. After treatment, 13/26 CSP patients and 11/24 FFP had greater than 50% normal protease activity. However, 66% of patients showed less than 10% inhibition of normal plasma protease activity indicating little, if any, inhibitor, in their plasma. Only 5 patients showed 100% inhibition of protease in mixing studies.

Conclusion: High levels of protease inhibitor do not always correlate with low platelet count or patient outcome. CSP did not show a significant advantage over FFP in this series.

O 40
Decreased Mortality Rate with Plasma Exchange in Patients with Class 1 HELLP Syndrome

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Erciyes University Faculty of Medicine, Department of Hematology, Department of Internal Medicine and Critical Care, Department of Gynecology and Obstetrics, Erciyes University, Kayseri, Turkey

Objective: Our purpose was to investigate the postpartum use of plasma exchange in patients with HELLP (hemolysis, elevated liver enzymes, and low platelets) syndrome with one or more risk factors.

Study Design: During two-year period (2000–2002) 23 patients with HELLP syndrome were treated post partum with single or multiple plasma exchange with fresh-frozen plasma (group I). All procedures were performed with Fresenius AS TEC 204 cell separator. Control group consisted of 26 patients with HELLP syndrome treated conservatively in between 1993 and 1999 (group II). All patients have had single- or multiple-risk factors (which were suggested by Martin et al.: platelet nadir < 50,000/µl, LDH level > 1,400 IU/L or AST level > 150 IU/L). Maternal complications and mortality rate were the main outcomes in this study.

Results: Median age, LDH level, platelet nadir and Hb level were not statistically different in two groups. One plasma volume corresponding to 40 ml/kg of body weight was exchanged daily until normal LDH level or normal platelet count were reached. Median five apheresis procedure (1–26) were performed. While maternal mortality rate was 23.1% (6/26) in group II, there was no death in group I. All deaths were in patients Class 1 with platelet nadir < 50,000/µl. The causes of mortality were infection, brain edema and ARDS. Mortality rate was significantly higher in patients with Class 1 (p<0.005) and all of group II patients (p<0.01). Dialysis requirement was not statistically significant in group I and group II (17.4% vs. 23%). Long term complications were observed in two group I patients and four group II patients.

Conclusion: Our study shows that plasma exchange reduces mortality rate in high risk patients with HELLP syndrome, particularly in Class 1 patients. Therefore, we suggest that plasma exchange should be added to standard therapy in patients with Class 1 HELLP syndrome and may be considered in other high risk group of patients.

L 43
The INTERCEPT Blood System for Platelets Provides Pathogen Inactivation while Maintaining Platelet Therapeutic Efficacy

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Cerus Corporation, Concord, California, USA

Background: INTERCEPT Blood System for platelets (plt) (IP), uses Helenix technology (amotosalen HCI [S-59] and UVA light) to inactivate a broad spectrum of bacteria, viruses and protozoa, as well as contaminating leukocytes. Nine clinical trials of IP were conducted. Recovery and lifespan in healthy subjects was shown to be within a therapeutically acceptable range. Prolonged template bleeding times in thrombocytopenic (tcp) patients (pts) were corrected with IP to a similar extent as with conventional plt (CP). Four more trials evaluated therapeutic efficacy and safety of IP in tcp pts requiring multiple plt transfusions (tx).

Methods: Two Phase 3 prospective, randomized, blinded trials compared IP processed with a clinical prototype device to CP. In SPRINT, 645 pts received Amicus apheresis IP or CP for up to 4 weeks. In euroSPRITE, 103 pts received tx with buffy coat IP or CP for up to 8 weeks. The 1st endpoint of SPRINT was 8% of pts with moderate (WHO Grade 2 bleeding); the 1st endpoint of euroSPRITE was 1-hour plt count increment (CI) and corrected CI (CCI).

Results: Demographics and baseline data were well-balanced in both studies.

<table>
<thead>
<tr>
<th></th>
<th>SPRINT</th>
<th>euroSPRITE</th>
</tr>
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<tbody>
<tr>
<td>N</td>
<td>318</td>
<td>327</td>
</tr>
<tr>
<td>% Grade 2 bleeding</td>
<td>58.5</td>
<td>57.5</td>
</tr>
<tr>
<td>% Grade 3 or 4 bleeding</td>
<td>4.1</td>
<td>NA</td>
</tr>
<tr>
<td>1-hr/24-hr CI (x10²/L)</td>
<td>21*13</td>
<td>34/22</td>
</tr>
<tr>
<td>1-hr/24-hr CCI (x10³)</td>
<td>11.1<em>16.7</em></td>
<td>16.0/10.1</td>
</tr>
<tr>
<td>Aplt dose (x10¹³)</td>
<td>4.6</td>
<td>4.0</td>
</tr>
<tr>
<td>Mean total plt dose (x10¹³)</td>
<td>29.4*</td>
<td>24.1</td>
</tr>
<tr>
<td>Mean no. plt tx</td>
<td>8.4*</td>
<td>6.2</td>
</tr>
<tr>
<td>Time between tx (d)</td>
<td>1.9*</td>
<td>2.4</td>
</tr>
<tr>
<td>Duration plt support (d)</td>
<td>11.5</td>
<td>10.5</td>
</tr>
<tr>
<td>Plt reaction (per tx)</td>
<td>5*</td>
<td>4</td>
</tr>
<tr>
<td>RBC tx</td>
<td>4.8</td>
<td>4.3</td>
</tr>
</tbody>
</table>

*p<0.05 compared to CP in same study.

By longitudinal regression analysis, 1-hr CI in euroSPRITE was not statistically different between IP and CP. Safety was comparable between IP and CP in both studies. No antibodies to S-59 neoantigens were confirmed. Two confirmatory trials in Europe with buffy coat and Amicus IP processed with the final device prototype confirmed these findings.

Conclusions: Therapeutic efficacy and safety of INTERCEPT platelets was demonstrated in a series of clinical trials in pts requiring multiple platelet transfusions. These results suggest that INTERCEPT platelets may be used whenever platelet transfusions are indicated.

O 44
Prevention of Graft-versus-Host Disease Using a Pathogen Reduction Process for Blood Products Using Riboflavin and UV Illumination

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Navigant Biotechnologies, 1215 Quail Street, Lakewood, CO 80215 USA

Objective: Transfusion associated Graft versus Host Disease (TA-GvHD) is a life threatening T cell mediated immune reaction in immunocompromised patients who receive blood products. Here we have evaluated the efficacy of a pathogen inactivation technology based on riboflavin and UV illumination to prevent TA-GvHD in a well characterized parent to F1 murine transfusion model.

Materials and Methods: B6AFl mice received an intravenous transfusion of either untreated B6AFl spleen cells (Group 1 – control Group), untreated A spleen cells (Group 2 – GvHD group), or A spleen cells treated with 50µM riboflavin, and illuminated with UV at a dose of 7.2 J/cm² (Group 3). The mice were observed and scored for clinical signs of TA-GvHD for 14 days. At termination, all mice were bled for hematological evaluation. The spleens were weighed and processed for analysis by flow cytometry. Liver, spleen skin and femurs were also collected for histopathology evaluation.

Results: Flow Cytometric Analysis

The percent of CD3+ T cells that were H-2K² positive was significantly lower in the GvHD group compared to both the control and treatment groups. This implies that there was less infiltration of the donor cells into the spleens of these animals compared to the GvHD group.

Splenomegaly
Both the spleen weights and the spleen to body weight ratio were significantly increased (p < 0.001) in the GvHD group (Group 2) compared to...
Pre-Transplant In Vitro Identification and In Vivo Monitoring of Donor-Derived Alloreactive T Cell Clones and Anti-Leukemia T-Cell Clones

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Objectives: Although graft-versus-host disease (GVHD) is usually associated with a graft-versus-leukemia (GVL) effect, GVCL can occur in the absence of clinical GVHD. A pronounced GVL effect following allogeneic hematopoietic stem cell transplantation (HSCT) has been documented in animal studies and by clinical observations. However, direct evidence that GVCL and GVHD are mediated by different clones of T cells is lacking.

Methods: Using irradiated peripheral blood mononuclear cells from patients in remission as target cells for GVH and irradiated leukemia cells from the same patient as target cells for GVL, we have demonstrated that donor-derived T cell clones when mediate GVH and GVL can be different. Primary mixed lymphocyte reactions (MLR) were performed using the recipient’s remission cells to obtain donor-derived GVH-specific T cell clones. In order to obtain donor-derived GVL-specific T-Cell clones, half of the cells in the primary MLR were treated with an anti-CD25 immunoconjugate to eliminate activated alloreactive T cells. The cells remaining after immunotoxin treatment were then incubated in a secondary MLR with the recipient’s leukemia cells. Activated CD4+ T cells from both the primary and secondary cultures were sorted on a FACS. mRNA was isolated from the sorted cells and a CDNA library was constructed. DNA sequences from the VDJ region encoding the T-cell receptor beta chain were determined.

Results: Ten patients with acute myeloid leukemia were analyzed. In one patient who relapsed after allogeneic HSCT, 3 recipient cell-specific and 2 leukemia cell-specific donor-derived CD4+ T cell clones were obtained. Both leukemia-specific clones were identical when pre-transplant or relapsing leukemia cells were used as targets in the MLR. One clone recognized both normal and leukemia cells. In a second patient two donor-derived GVH-specific clones and one GVL-specific clone were identified. Remaining 8 patients had HLA-mismatched healthy donors only available for in vitro experiments. In all 8 cases the presence of different donor-derived GVH- and/or GVL-specific T-cell clones was clearly demonstrated.

Conclusions: Our recent data demonstrate direct evidence that GVH- and GVL-specific T-cell clones can be separated in vitro and individually monitored in vivo. This strategy is being used to selectively deplete GVH-specific and expand GVL-specific donor-derived T cells prior to donor lymphocyte infusion.

O 46 Controlled-Rate vs. Uncontrolled-Rate Freezing of Platelets – What Is Better?

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Objective: In an attempt to establish the optimum method for platelet cryopreservation, much work has been carried out. We described our experience with cryopreservation of platelet concentrate derived from buffy-coat (PC-BC) using six different protocols for platelet cryopreservation.

Methods: PC-BC units were examined before and after cryopreservation – using uncontrolled-rate and original controlled-rate freezing (with compensation of released heat of fusion) and 6% or 10% dimethyl sulfoxide (DMSO) and 5% DMSO with hydroxethyl starch (HES), as cryoprotective agents. The platelet count was determined by a flow cytometer Technicon H-3. Morphology of platelets was examined using phase-contrast microscope technique (Polyvar). Four type of platelets were characterized (discoid = 4, spheres = 2, dendrites = 1, balloons = 0 points) to calculate the platelet morphological score (PMS). Platelet ultrastructure was investigated by electronmicroscope (Philips 201C). Cell functions were estimated using tests of hypotonic shock response (HSR) and platelet aggregation with ADP.

Results: Post-thaw platelet recovery in PC-BC groups cryopreserved with 6% DMSO was 91.0 ± 5.5 (controlled-rate freezing) and 85.9 ± 6.5% (uncontrolled-rate freezing), respectively (p < 0.01). The application of controlled-rate freezing resulted in better PMS-recovery (81.8 ± 2.8%) than uncontrolled-rate freezing (75.7 ± 3.9%). The percentage of discoid platelets were higher in controlled-rate setting (57.9 ± 2.6%) than in uncontrolled-rate setting (51.6 ± 3.7%), too. Ultrastructural investigation showed the high frequency of platelets with degranulation, and appearance of pseudopodes and vacuoles when HES and DMSO were used. Finally, the best answer in HSR-test were obtained in cryopreserved with controlled-rate rate of freezing and 6% DMSO (68.0 ± 23.2%). The aggregation in the same group compare to control group was 44.8 ± 13.2%.

Conclusion: Our original cryopreservation procedure using controled-rate of freezing (with compensation of released heat of fusion) and 6% DMSO resulted in highest percent of viable platelets vs. other used freezing methods.

O 50 Observations on Protein A Immunoadsorption as an Adjunctive Therapy for Acquired Thrombotic Thrombocytopenic Purpura

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Purpose: According to preliminary results, Protein A immunoadsorption (PAIA) should be useful in the treatment strategy of several severe autoantibody mediated haematological disorders. We summarize the data on our first patient (P) with acquired thrombotic thrombocytopenic purpura (TTP).

Method: In TTP plasma exchange (PE) with fresh frozen plasma (FFP) replacement is standard management, removing circulating inhibitors and providing exogenous ADAMTS13 activity. We investigated, whether PAIA can induce a faster clinical response by removing circulating inhibitors more efficiently than PE and thereby restoring enzymatic enzyme activity more rapidly.

Results: For PAIA and PE we utilize a cell separator (Spectra, Gambro BCT, USA). In PAIA the secondary system consists of staphylococcal Protein A- columns and a plasma flow monitor (Immunosorba and Citem 10, Fresenius, Germany). In every single PE (PAIA) 1.0–1.5 (2.0–2.5) plasma volumes of the P were replaced with FFP (processed). We started with 3 daily PE and switched to PAIA on day 4–6. Then PAIA was discontinued and PE resumed for another 10 days. Further treatment consisted of Methylprednisolone 125 mg/day i.v.

Results:

<table>
<thead>
<tr>
<th>Day</th>
<th>PE/PAIA</th>
<th>Platelets (G/L)</th>
<th>LDH (U/L)</th>
<th>ADAMTS 13 Activity (%)</th>
<th>ADAMTS 13 Inhibitor (BU)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>PE</td>
<td>7</td>
<td>3,224</td>
<td>&lt;3%</td>
<td>ca. 3</td>
</tr>
<tr>
<td>4</td>
<td>PAIA</td>
<td>70</td>
<td>606</td>
<td>5%</td>
<td>&lt;1</td>
</tr>
<tr>
<td>7</td>
<td>PE</td>
<td>17</td>
<td>2,571</td>
<td>&lt;3%</td>
<td>&lt;1</td>
</tr>
<tr>
<td>14</td>
<td>PE</td>
<td>108</td>
<td>495</td>
<td>30%</td>
<td>&lt;= 1</td>
</tr>
</tbody>
</table>

Initial 3 PE transiently improved platelet count and LDH. This paralleled with normalized ADAMTS13 activity (prior/after 3rd PE: 5% / > 50%) and lowered the inhibitor titer. After switching to PAIA disease activity deteriorated (see table). ADAMTS13 activity returned to < 5% within < 20 h although the inhibitor titer remained virtually undetectable during PAIA. Resumption of PE again led to an improvement of clinical
and laboratory parameters. Conclusions: PAIA neither improves disease activity nor restores ADAMTS13 activity, although circulating ADAMTS13 inhibitors are virtually completely removed. Residual traces of the inhibitor or its re-entry from extravascular sites seem to be sufficient for completely inhibiting newly synthesized endogenous ADAMTS13 and/or the enzyme is produced insufficiently. For future applications to acquired TTP, PAIA should be combined with infusion of exogenous ADAMTS13.

O 51 Further Experience with Protein A Immunoadsorption in Autoimmune Haemophilia

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Purpose: Extracorporeal immunoadsorption to staphylococcal Protein A (PAIA) is a powerful tool in reducing circulating IgG and immune complexes. According to preliminary results, its implementation in the treatment strategy of several severe autoantibody mediated haematological disorders (e.g. catastrophic antiphospholipid-syndrome and thrombotic thrombocytopenic purpura) should be useful. Here we summarize our data on 6 homozygous haemophilia patients (PS) with factor VIII autoantibodies (FVIII-auto-AB; FVIII-auto-AB) which is one of the most promising indications.

Methods: We use a combination of PAIA, cyclophosphamide (CY), corticosteroids (CST) and immunoglobulins i.v. as induction cycles (IC). These are repeated every 3–4 weeks until consistent lowering of the inhibitor or its re-entry from extravascular sites seem to be sufficient for completely inhibiting newly synthesized endogenous ADAMTS13 and/or the enzyme is produced insufficiently. For future applications to acquired TTP, PAIA should be combined with infusion of exogenous ADAMTS13.

Results: The 8 patients who suffered from CAD were 16 to 38 years old (m = 29.8 ± 8.8 years). They started LDL apheresis at 5 to 33 years old (m = 17.8 ± 8.5 years) 1/18 patients had 1 additional risk factor (smoking). 5/8 had clinical and angiographic regression of CAD after onset of LDL apheresis. 1 patient, 38 years old, died from massive myocardial infarction, 18 years after onset of LA (he didn’t take any lipid-lowering drug).

Background and Hypothesis: The DALI system is a highly efficacious method for removing LDL in patients with hypercholesterolemia. However, some patients develop side-effects and are intolerant to this valuable mode of therapy. Among the serious adverse effects is a syndrome of faecal flushing, urticaria, and shortness of breath. This was observed in 6/22 of the patients undergoing DALI treatment in our hospital during the past 3 years (Durst et al., IMAJ 4:677, 2002). This syndrome was always noted in the initial phase of therapy (during the first 2–3 sessions). It was suspected that this syndrome was caused but the high concentration of citrate which is released from the column after priming and abruptly enters the patient’s circulation. Since this parameter was set automatically and was not under the user’s control, we could not test our hypothesis. This syndrome recurred in the 3 patients who agreed to be rechallenged under the same treatment conditions. Therefore, all patients experiencing this syndrome were discontinued from DALI treatment and were treated using other methods.

Objectives: To develop an alteration of the DALI system which would A- not cause this adverse reaction and B- elucidate the etiology of this adverse effect.

Methods: Because of the awareness of this problem, which had also been reported by other centers, the manufacturer (Fresenius) developed a modification of the system which uses half of the amount of citrate during low-citrate program, with no change in the column or other components of the system, these patients were able to undergo 6 DALI treatment each with absolutely no adverse effects. They are continuing DALI treatment as their regular LDL-lowering treatment modality. Two additional (sensitive but not frankly allergic) patients who had been treated with the high-citrate DALI system were switched to the low-citrate system. They reported feeling subjectively much better after low-citrate DALI sessions.
Conclusion: We conclude that the flushing/dyspnea seen using DALI LDL apheresis system is indeed a function of high citrate in the return line after priming and it can be successfully prevented using a low-citrate modification of the program.

L 62
Conversion of Platelet Production Method: From Buffy Coat Platelets to Single Donor Platelets – a Feasibility Study
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Objective: Since variant Creutzfeldt-Jakob (vCJD) became a potential threat for the blood supply the Dutch medical advisory board has advised Sanquin to take measures to minimize donor exposure. One of the recommendations was to transfuse only single donor platelets (SDP). Sanquin promoted a pilot study to evaluate technical, logistical and economical feasibility of thrombocytapheresis (TPH). To enhance economical feasibility it was decided to target double platelet products (DPP) only.

Donors and Methods: All A- and O- plasmapheresis donors (PPH) with platelet pre-count of ≥250,000 and weight >70 kg, who showed previous willingness to become TPH donors, were invited to donate. 100 procedures were performed on Trima™ Accel™ (Gambro BCT). All collected SDP were tested in the QC lab on platelet yield, residual leukocytes, volume, pH, swirl and sterility (bacterial screening). Plasma products were tested on volume and cell counts. Targeted procedure time was 70 min with an exception till 90 min.

Results and Discussion: 58% of participating donors met all the selection criteria. 86.2% of the procedures resulted in a DPP. Mean PLT Yield was 540 ± 99 × 10^11. WBC content complied with European guidelines. Average procedure time was 64.9 ± 69. 60 of the 100 procedures took at most 70 min, however, 23 procedures took between 70 and 90 min. Simulations on volume, pH, swirl and sterility (bacterial screening). Plasma products were tested on volume and cell counts. Targeted procedure time was 70 min with an exception till 90 min.

Conclusion: Conversion of buffy coat platelets to single donor platelets appears feasible. Good donor selection and the high collection performance assure high split rates and relatively short procedure times. It further showed that the centre would be auto sufficient (2100 SDP annually) if 203 donors would donate 6 times a year.

O 65
The Content of the LRS Chamber Provides a New Quality Tool for Characterization of the Donor Platelet Profile
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Objectives: to compare the relative Platelet (Plt) indices and aggregation states of the single dose platelet-derived from Trima with those obtained from the LRS chamber (LRS) and relate them with programmed yields.

Material and Methods: Immediately at the end of collection (n = 25), the LRS was disconnected from the system and the content removed according to a standard procedure. For a quantitative assessment of Plt aggregation and its response to edta, paired samples of edta and citrate were counted with a Cell Dyn 3500 and calculated for dPlt (Pltedta – Pltcitrate) and dMPV (MPvedta – MPvcitrate). The same approach was used for the products.

Results: on d0 there was a good correlation between donor and product MPV (8.4 versus 8.6; r = 0.84). Mean values of LRS MPV and dMPV (10.8 and 1.6, respectively) were suggestive of the presence of larger aggregates with different response to edta, as compared to those obtained from the product on d0 (1.01), d1 (1.2) and d2 (1.05). The products on d0 also showed a higher dPlt (mean = 222) but aggregates were reversible, as demonstrated by the fall in dPlt to 31 on d1 and to 15 on d2. The ratio between programmed/obtained yields varied from 0.8 to 1.1, except 2 donors who showed a donor-specific pattern. 4 donors had a Plt counting in LRS >2200 × 10^12/µL. The first donor presented several alarms during collection, requiring revision of patient. Yield ratio was 1.6 (3.2/2) and the ratios in the 2 subsequent collections were 1.4 and 2.5. Second donor had a ratio of 4.4 (3.2/0.8). LRS was full of macroscopic aggregates. A similar ratio had been obtained in a previous collection (3.2/0.7). Third donor presented a ratio of 0.9 (4/4.7) in spite of a LRS Plt of 2,534 × 10^12/µL. He was included in a double doses programme later on. No unusual pattern was found on the fourth donor.

Conclusions: The product Plt indices were found to be donor-specific, the larger Plt being retained in LRS chamber. The simultaneous analysis of products and LRS content provide useful information not only for the characterisation of donor-related phenomena, but also help in the identification of potential shortcomings in the machine performance allowing for remedial action taken on evidence based data.

O 66
The Quality of Intercept™ Treated Amicus™ Apheresis Platelets as Compared to Untreated Platelets
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Objective: We compared the quality of Intercept™ treated and untreated apheresis platelet concentrates (PC) and evaluated the in vitro effects of storage.

Materials and Methods: 20 double dose PCs have been collected with the Amicus™ and split immediately after apheresis. The platelet additive solution factor was set to 0.57% leading to an InterSol™ fraction of median 60% (range 53–65). 4 PCs had to be discarded because of visible aggregates. The half of each PC underwent Intercept™ treatment the day after production, the other half was left untreated.

Results: The platelet yield of the double dose PC was 7.18 ± 10^11 (5.35–8.07), and the amount of rWBC was 0.03 ± 10^6 (<0.02–0.45). Platelet loss after 7% (3–14).

<table>
<thead>
<tr>
<th>Day 2 before treatment</th>
<th>Intercept™ treated PCs</th>
<th>Untreated PCs</th>
</tr>
</thead>
<tbody>
<tr>
<td>6.65 (0.74)</td>
<td>6.62 (0.74)*</td>
<td>6.75 (0.67)</td>
</tr>
<tr>
<td>pH (22 °C)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>7.03 (0.09)</td>
<td>6.77 (0.11)*</td>
<td>6.94 (0.13)*</td>
</tr>
<tr>
<td>Glucose (mg/dL)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>115 (19.9)</td>
<td>113 (17.5)*</td>
<td>51 (25.1)*</td>
</tr>
<tr>
<td>Lactate (mmol/L)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>4.5 (1.2)</td>
<td>4.7 (2.0)*</td>
<td>11.3 (1.9)*</td>
</tr>
<tr>
<td>LDH (U/L)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>115 (56)</td>
<td>150 (58)*</td>
<td>205 (144)*</td>
</tr>
<tr>
<td>CD62p (%)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>14.5 (3.1)</td>
<td>16.9 (4.0)*</td>
<td>13.3 (3.7)*</td>
</tr>
<tr>
<td>HSR (%)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>59 (19)</td>
<td>61 (17)</td>
<td>56 (14)</td>
</tr>
</tbody>
</table>

Results are given in median and (standard deviation). * = p < 0.02 versus previous day. * = p < 0.02 versus treated group same day.

Conclusion: The Intercept™ treatment causes a platelet loss of approx. 7% which has to be considered for reaching a satisfying platelet dose for transfusion. pH levels are significantly reduced after Intercept™ treatment due to the addition of 15 mL Amotosalen™ (pH = 4.5). The continuing effect of the Intercept™ treatment is reflected by the further decrease of the pH level, higher platelet activation, and larger platelet volumes at the end of storage as compared to untreated platelets. Platelets’
metabolism and their ability to respond to hypotonic shock are not affected by the Intercept® treatment. The formation of visible aggregates which did not dissolve during storage over night maybe due to the separation technology of the Amicus® and/or to the addition of InterSolTM. Platelets suspended in InterSol™ should not be stored > 5 days because glucose and pH levels are very low at day 7 due to the contents of 60% InterSol™.

O 67
Implementation of Concurrent Red Blood Cell and Platelet Collection by Apheresis
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Background and Objectives: New technological developments make it possible to collect red blood cells (RBCs) by apheresis which provides standardised products and has the potential for improved RBC quality. Concurrent collection of RBCs and platelets (PLTs) allows for an increase of the blood supply and reduces costs of laboratory tests. The present study analyses the number of concurrently collected RBC units in plateletpheresis donors and the reasons why donors were deferred from multicomponent collection.

Material and Methods: Donors fulfilling inclusion criteria for multicomponent donation underwent concurrent collection of RBCs and PLTs with the single needle procedure of the Amicus blood cell separator. The haemoglobin value prior to RBC collection and of the follow up donation as well as the reasons for deferral were retrospectively evaluated for a period of one year.

Results: A total of 404 RBC units were concurrently collected with PLTs. An average of 1.8 RBC units was collected from each donor. The baseline haemoglobin value was almost equal for the first (n = 221), the second (n = 117), the third (n = 54) and the fourth donation (n = 12). Concurrent PLT and RBC collections were well tolerated by most donors. An RBC unit was not collected in 190 aphereses. 39 donors (20.5%) were not accepted for RBC collection due to a donation interval of less than three months. Haematoma and blood flow problems occurred in 36 (18.9%) recipients. A total of 190 aphereses failed due to technical reasons. Donors fulfilling inclusion criteria for multicomponent donation underwent concurrent collection of RBCs and PLTs with the single needle procedure of the Amicus blood cell separator. The haemoglobin value prior to RBC collection and of the follow up donation as well as the reasons for deferral were retrospectively evaluated for a period of one year.

Conclusion: RBC supply was increased by the implementation of concurrent RBC and PLT collection. The donor eligibility for the procedure has to be taken into account and the logistics of RBC processing (filtration) should be optimized for a further increase of blood supply.

O 69
Data Management in Preparative Hemapheresis
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Objective: Systems that ensure quality in Blood Centres are submitted to more and more stringent requirements these last years. We studied the benefits of recording hemapheresis data from a technical point of view.

Study Design: 4 combinations of data management systems and hemapheresis devices were compared:
1. Apheresismaster/ComTec
2. HaemoNet/MCS+
3. Vista/Trima
4. Ami-Print/Amicus

Parameters that can influence the quality of the platelet concentrates can be divided into 2 groups:
1. Procedure related conditions such as: ACD-A consumption (inlet; AC ratio, ACD-A in the product in relation to the expiry and storage of the product), continuity of the blood flow, procedure time (regeneration of platelets from the spleen), alarms, cell concentration, tubing set (length of the set determines the residual set volume, blood loss tracking), ...
2. Donor related conditions: pre-hematocrit, platelet count, mean platelet volume, white blood cell precount, ...

The donor and procedure data recorded in the 4 data management systems was compared. We also evaluated the influence of registration and assessment of procedure data on the final quality of the platelet concentrates. Also the obtained messages and alarms were listed. Special attention was given to the statistics these software programs offer, together with the databases used. In this aspect is could be stated that the databases of these software programs use, are composed of two interconnected parts. The first one is a donor database containing all donor related info such as name, address, gender, height, weight, previous platelet and hematocrit counts, etc. Databases (like Vista) keep also track of the blood loss of the donor. The second part of the database is the procedure database.

Summary: Although there exist some similarities between the 4 software/vice combinations, we could demonstrate clear differences between the systems especially in the field of QC, AC management and blood loss tracking.

P 3
Comparison of the LDP C5 and LDP C2 Protocols for Platelet Concentrates Preparation on Haemonetics MCS+ Separator
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Objective: The purpose of this study was to compare the quality of platelet concentrates collected with two different software versions on the same cell separator.

Methods: 23 platelet apheresis procedures were performed using Haemonetics MCS+ separator, protocol LDP – new revision C5. The results were compared with 20 procedures performed on the same MCS+, protocol LDP – revision C2. 994CF-E sets were used in all apheresis. Donor characteristics, platelet yield, platelet volume, residual WBC count, collection efficiency (CE), and side effects were recorded and analysed.

Results: Donor data did not differ significantly. In the C5 group there was slightly higher CE (60.4 ± 6.8%, range 48.8–70.1%, median 61.4%) than in C2 group (59.8 ± 4.5%, range 51.5–70.2%, median 59.5%, p < 0.05). Calculated platelet yield per cycle was 62.4 ± 11.5 x 10^9, range 48.7–97.5 x 10^9, median 61.4 x 10^9 in the C2 group and 67.1 ± 10.5 x 10^9, range 42.0–88.5 x 10^9, median 60.2 x 10^9, in the C2 group (p < 0.05). Calculated platelet yield per hour was (C5): 2.96 ± 0.57 x 10^11, range 1.95–3.76 x 10^11, median 3.03 x 10^11 and (C2): 2.94 ± 0.49 x 10^11, range 2.21–4.01 x 10^11, median 2.94 x 10^11 (p = 0.05). No side effects were observed. The WBC contamination (Nageotte chamber) was lower than 1 x 10^11 in all platelet concentrates.

Conclusions: Results of platelet concentrates collected using the protocol LDP-C5 revision and the protocol LDP-C2 revision were comparable. There were no statistically significant differences comparing procedures using these two revisions.

P 4
Results of a Productivity Comparison of Trima® Accel™ with Haemonetics-MCS+ in Platelet Collections
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Background: To compare the productivity of two plateletpheresis cell separators (Trima® Accel™ and Haemonetics MCS+). Relevant donor parameters (age, sex, total blood volume, hematocrit, platelet count and mean platelet volume), product parameters (volume, predicted yield and real yield) and procedure parameters (time until update of donor parameters, volume of blood processed and procedure time, incidences) were collected. For statistical analysis chi-square and t-test evaluations were used. For correlation use was made of the regression model and correlation factor r of Pearson.

Results: Donor characteristics were similar in both groups. With regard to procedures Trima® Accel™ V5.0 and 173 procedures with Haemonetics-MCS+. Relevant donor parameters (age, sex, total blood volume, hematocrit, platelet count and mean platelet volume), product parameters (volume, predicted yield and real yield) and procedure parameters (time until update of donor parameters, volume of blood processed and procedure time, incidences) were collected. For statistical analysis chi-square and t-test evaluations were used. For correlation use was made of the regression model and correlation factor r of Pearson.

Results: Donor characteristics were similar in both groups. With regard to procedures Trima® Accel™ V5.0 and 173 procedures with Haemonetics-MCS+. Relevant donor parameters (age, sex, total blood volume, hematocrit, platelet count and mean platelet volume), product parameters (volume, predicted yield and real yield) and procedure parameters (time until update of donor parameters, volume of blood processed and procedure time, incidences) were collected. For statistical analysis chi-square and t-test evaluations were used. For correlation use was made of the regression model and correlation factor r of Pearson.
good for both cell separators albeit that the correlation was higher with Trima<sup>®</sup> Accel<sup>™</sup> (r = 0.87 vs. MCS+ r = 0.78). Approximately 40% of the donors showed a variation above 10% between predicted and targeted yield. This was not caused by the time required to update the donor parameters after the start of the procedure (mean time = 11 min, percentiles25 = 8 y percentiles75 = 13). A total of 12 infinations were observed with Trima<sup>®</sup> Accel<sup>™</sup> V5.0 (3.3%) and 4 with MCS+ (2.3%). This difference was statistically not significant (p = 0.518). There were 2 vasovagal reaction that required early termination of the procedure (1 on each platform).

**Conclusions:** Absolute yield as well as yield per minute were superior with Trima<sup>®</sup> Accel<sup>™</sup> V5.0. Both platforms are well tolerated and achieve adequate platelet product yields.

**P 5**

**Collection of Plasma-Reduced Platelet Concentrates on Trima<sup>®</sup> Accel<sup>™</sup>, Compared to Collection of Standard Concentrates on Trima Version 4**

R. Verheyden, L. Steensens

**Blood Transfusion Center Vlaams Brabant-Limburg, Leuven, Belgium**

**Background:** In January 2003 one Trima version 4 (v4) automated blood collection system was upgraded to TRIMA ACCEL. Plasmareduced platelet concentrates (PPC) collected on TRIMA ACCEL were compared with normal platelet concentrates (NPC) collected on v4 in terms of yield, productivity and cellular contamination.

**Methods:** Per donor each TRIMA ACCEL procedure (n = 74; all different donors) was compared to the next v4 procedure of the same donor. Collection concentration for TRIMA ACCEL was 4,100 x 10<sup>9</sup> for Trima v4. A prediction-dependent volume of T-sol was added to the PPC within 0.5 hour after collection to reduce the concentration to normal levels. This way, after addition of T-sol PPC had a plasma carryover of approximately 37%.

**Results:** See table 1 above.

**Table 1.**

<table>
<thead>
<tr>
<th>Predicted yield</th>
<th>Predicted yield</th>
<th>Predicted yield</th>
<th>Efficiency (%)</th>
<th>CR (plt/min)</th>
<th>Procedure failures</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.8 x 10&lt;sup&gt;11&lt;/sup&gt;</td>
<td>3.2 x 10&lt;sup&gt;11&lt;/sup&gt;</td>
<td>4 x 10&lt;sup&gt;11&lt;/sup&gt;</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Tr1 n = 50</td>
<td>n = 58</td>
<td>n = 10</td>
<td>n = 99</td>
<td>n = 121</td>
<td>n = 9</td>
</tr>
<tr>
<td>2.79 ± 0.49</td>
<td>3.27 ± 0.46</td>
<td>4.13 ± 0.66</td>
<td>59.51 ± 9.8</td>
<td>6.6 ± 0.01</td>
<td>7%</td>
</tr>
<tr>
<td>Tr2 n = 47</td>
<td>n = 55</td>
<td>n = 4</td>
<td>n = 85</td>
<td>n = 109</td>
<td>n = 3</td>
</tr>
<tr>
<td>2.83 ± 0.38</td>
<td>3.27 ± 0.46</td>
<td>4.32 ± 0.45</td>
<td>60 ± 7.1</td>
<td>6.6 ± 0.0</td>
<td>3%</td>
</tr>
</tbody>
</table>

**Table 2.**

<table>
<thead>
<tr>
<th>pH Measurement</th>
<th>Tr1 d&lt;sub&gt;0&lt;/sub&gt;</th>
<th>Tr1 d&lt;sub&gt;1&lt;/sub&gt;</th>
<th>Tr1 d&lt;sub&gt;5&lt;/sub&gt;</th>
<th>Tr2 d&lt;sub&gt;0&lt;/sub&gt;</th>
<th>Tr2 d&lt;sub&gt;1&lt;/sub&gt;</th>
<th>Tr2 d&lt;sub&gt;5&lt;/sub&gt;</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 h</td>
<td>n = 10</td>
<td>n = 11</td>
<td>n = 11</td>
<td>n = 8</td>
<td>n = 8</td>
<td>n = 8</td>
</tr>
<tr>
<td>7.21 ± 0.08</td>
<td>7.5 ± 0.12</td>
<td>7.38 ± 0.12</td>
<td>7.19 ± 0.02</td>
<td>7.48 ± 0.04</td>
<td>7.43 ± 0.1</td>
<td></td>
</tr>
<tr>
<td>3 h</td>
<td>n = 6</td>
<td>n = 8</td>
<td>n = 8</td>
<td>n = 8</td>
<td>n = 8</td>
<td>n = 8</td>
</tr>
<tr>
<td>7.26 ± 0.07</td>
<td>7.4 ± 0.03</td>
<td>7.38 ± 0.07</td>
<td>7.26 ± 0.06</td>
<td>7.44 ± 0.08</td>
<td>7.42 ± 0.19</td>
<td></td>
</tr>
</tbody>
</table>

**Conclusions:** pH remained at the upper limit of specification. The conformance with predefined requirements allowed us to consider the system validated. We concluded that products obtained by Trima are acceptable and provide safe leucodepleted concentrates for clinical use.

**P 6**

**Trima Validation for Single Apheresis Platelet Products**

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**Objectives:** to validate the collection and storage characteristics of single dose platelets (plt) obtained by 2 Trimas (Tr1 and Tr2) against CE guidelines.

**Material and Methods:** 224 platelepheresis products, obtained from healthy donors (plt > 200,000/μL) were evaluated. Collection was performed according to the manufacturer instructions. The target yields were 2.8, 3.2 and 4x10<sup>11</sup> platelets/unit. Regarding pH measurement, on an initial phase 44 products with 1 hour of resting period (rp) were evaluated on day 5 (d5). On a subsequent phase, evaluation was performed on a total of 30 products and included measurements on d1 before shaking, d2 and d5. The rp was 3 and 8 hours for 19 and 11 products, respectively. A Cell Dyn 3500 counter, BD Facsicalibur flow cytometer, Chiron 340 meter and Bact Alert system were used for plt counting, leucocyte evaluation, pH measurement and bacteriological screening. The overall performance of both machines was assessed by collection rate (CR) and process efficiency.

**Results:** The obtained values of CR, process efficiency and yields are summarized in table 1. There was a correlation of 0.91 between programmed and final volumes. Leucoreduction specification was attained on a 100% basis, as none contained more than 1 x 10<sup>6</sup> leucocytes. 212 products contained more than 2 x 10<sup>11</sup> platelets/unit, which corresponded to a 95% degree of conformance to the CE guidelines. No bacterial contamination was found. No effects on pH were obtained after enlarging the rp (table 2).

**P 7**

**Quality Control in Apheresis Platelet Concentrates: Various WBC Reduction Technologies Lead to Different Leukocyte Depletion**


Clinical Blood Group Serology and Transfusion Medicine, Vienna University Hospital, Austria

**Objective:** The guidelines of the Council of Europe require the amount of rWBCs in a leucocyte depleted platelet concentrate (PC) < 1 x 10<sup>6</sup>. WBC reduction of apheresis PCs can be performed by preparative technologies (fluidized particle bed technology, elutration principle, interface detector) or by filtration.
Material and Methods: At least four PCs per month of each cell separator are randomly assigned to our QC laboratory. Measurement of rWBCs is performed with the Leuco-Count kit (B and D). If a PC contains >1×10^9 rWBCs, the next two PCs produced by this cell separator also undergo QC. If a second PC > 1×10^9 rWBCs is detected, the cell separator is eliminated from production till a technical service has been performed. The first PC after reparation is controlled again.

Results: A total number of 4,965 PCs have been produced in 2002. QC was performed in 962 PCs (19.4%).

<table>
<thead>
<tr>
<th>Cell separator</th>
<th># PCs</th>
<th># QCs</th>
<th>rWBC (&gt;10^9)</th>
<th># &gt; 1×10^9</th>
<th>median (range)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Amicus SN</td>
<td>803</td>
<td>319</td>
<td>0.04 (&lt;0.02–9.99)</td>
<td>34 (11%)</td>
<td></td>
</tr>
<tr>
<td>Cobe Spectra DN</td>
<td>411</td>
<td>70</td>
<td>0.04 (&lt;0.02–29.3)</td>
<td>1 (1.5%)</td>
<td></td>
</tr>
<tr>
<td>Cobe Trima</td>
<td>425</td>
<td>53</td>
<td>0.04 (&lt;0.02–6.9)</td>
<td>2 (4%)</td>
<td></td>
</tr>
<tr>
<td>ComTec</td>
<td>295</td>
<td>97</td>
<td>0.03 (&lt;0.02–3.9)</td>
<td>11 (11%)</td>
<td></td>
</tr>
<tr>
<td>MCS+</td>
<td>3,030</td>
<td>423</td>
<td>&lt;0.02 (&lt;0.02–364)</td>
<td>38 (9%)</td>
<td></td>
</tr>
</tbody>
</table>

Conclusion: The interface detector which is responsible for WBC reduction with the Amicus (Baxter) and the ComTec (Fresenius) is very sensitive and high rWBCs are found in 11% of the PCs. PCs from the MCS+ (Haemonetics) show the lowest leukocytone contamination in median, but the incidence of filtration failures is 9%. The participle bed technology used with the Cobe has constantly good rWBC results. In some cases failed WBC reduction is donor related: these donors are identified, and they are assigned to another type of cell separator.

P 8 Clinical Efficacy of Plasma-Reduced (PR) Platelet Concentrates from Multi-Component (MC) Collection: A Prospective Study

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Servizio Trasfusionale, Ospedale San Gerardo, Monza and *Clinica Pediatria-CTMO,Univ. Milano-Bicocca

Background: PR-MC platelet collection has been recently introduced in several blood collection units, with both the aims of saving plasma for industry and to possibly reduce the incidence of plasma-related side-effects (SE) in patients receiving multiple PLT transfusions (TX). We performed a single center non-randomized prospective study to compare PR-MC platelet units to standard random PLTs and to platelethropheresis units.

Patients and methods: Onco-haematological pediatric pts who were given prophylactic PLT TX for PLT values <20×10^9/L or for bleeding episodes, entered the study. They received PR-MC PLT units obtained by using Cobe TRIMA LRS device, random PLTs (R-PLT) or platelethropheresis obtained by using Cobe Spectra LRS Turbo device (PLT-P), depending on the product availability and their body weight, with the aim to infuse at least 0.5×10^11/dl PLTs. ATP, ADP and AMP immediately after collection, postthawing, after 24 h and 1 week. The tolernance was evaluated in 66 pts received a total of 221 PLT TXs: 92 pts were given R-PLT, 58 PLT-P and 71 PR-MC PLT. The 3 groups did not differ in their pre-TX PLT count, whilst those who were given R-PLT had a lower bw compared to PLT-P and PR-MC PLT groups: 30 ± 19 Kg vs. 41 ± 19 Kg and 43 ± 21 Kg, respectively (p < 0.05). As to PLT TX clinical efficacy, +1 h CCI was 15.7 ± 9.11 in 15 and 13 + 9 (p = n) and +24 h CCI was 9 ± 8.7 ± 13 and 9.4 ± 7.9 (p = n) in the 3 groups. PLT TX-related side effects were observed in 6 out of 92 R-PLT TXs (6.5%) and in 2 out of 71 PR-MC PLT TX (2.8%), no SE was observed in pts who were given PLT-P units.

Conclusions: Pts who were given PR-MC PLT units showed a satisfactory post-TX CCI and, similarly to those who received PLT-P units, a very low incidence of plasma-protein related post-TX SE, when compared to pts who were given R-PLT units. PR-MC PLT units proved their safety and clinical efficacy in children from an onco-haematological setting.

P 9 Double Red Blood Cell Apheresis and Clinical Applications

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1Antalya State Hospital, Thalassemia Center and Blood Center-Antalya, 2Suleyman Demirel University, Blood Center -Isparta, Turkey

Objective: Erythroapheresis has two types; donor erythroapheresis including single, double and neocyte and therapeutical erythroapheresis including erythrophycytoreduction and erythrocyte exchange. Double Red Blood Cell Apheresis (DRBCA) has many advantages such as maintaining standard red cell volume and hematorcit content, decreasing the risk of alloimmunization and blood-borne infections, increasing red blood cell quality and efficacy for patients. Although it seems high cost, the analysis of cost-effect shows normal cost because only one pretransfusion test is performed for double unit red blood cell unit, alloimmunization risk is decreased by using pre storage log 4 leukocyte filters.

Material and Methods: All 605 voluntary unpaid blood donors were carried out a Double RBC Apheresis by Haemonetics MCS+ including an inline Pall filter at Antalya State Hospital Blood Center between August 2000 and December 2002.

Results: The mean age of donors (604 male, 1 female) were 33 ± 8 years, their mean pre donation Hb levels: 15.4 ± 1 g/dl, hematocrit levels: 45.7 ± 2%, WBC: 8.1 ± 5 ×10^9/L and platelet: 251.2 ± 56×10^9/dl. The tolerance of all donors were very good and had no adverse reactions. Their blood group distribution were as follows: A+ in 239 (39.5%), O+ in 221 (36.5%), B+ in 71 (11.7%), AB+ in 43 (7.1%), A− in 20 (3.3%), O− in 5 (8.3%) and B− in 3 (0.5%) patients. 120 of 1,210 RBC units (9.9%) for patients with thalassemia major, 128 units (10.5%) for the patients with malignancy and 962 units (79.5%) for open heart surgery patients at the department of cardiovascular surgery were used.

Conclusion: Double RBC Apheresis is safe and reliable for donors and maintains high quality and safe hemotherapy for all patients.

P 10 Stability of Thawed Packed Red Blood Cells Using the ACP 215® of Haemonetics for Freezing and Thawing

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Clinical Department for Transfusion Medicine. * Department of Cardiothoracic Surgery, University Hospital, Vienna, Austria

Background: In the freeze and thaw technology of packed red blood cells (PRBCs) commonly open systems are used. Therefore the shelf life of thawed PRBCs is limited to 24 hours. In this study we evaluated an automatic and functionally closed system for both the glycerolization and deglycerolization process which allows a postthawing storage of 7 days.

Material and Methods: Seventeen PRBCs were collected from healthy male donors using the MCS+ device. All donors met the national and European guidelines for eligibility for cytapheresis donors and gave written informed consent to participate in the study. PRBCs were glycerolized immediately after donation (without adding SAGM) using the high glyceral method. Glycerolization was done with the ACP® device (Haemonetics) at a final concentration of glycerol of 37%. These PRBCs were stored at ~80° for 14 days. Quality was assessed by the measuring of cell counts, free hemoglobin (fHb), K+ LDH, pH, lactate, glucose and intracellular ATP, ADP and AMP immediately after collection, postthawing, after 24 hours, on day 4 and 7.

Results: Rate of hemolysis as determined by fHb, LDH, Hct as well as the increase of K+ content in the supernatant, lactate and glucose remained within the ranges of conventionally stored PRBCs during their shelf life. Over the storage period pH remained stable above 6.5 and the recovery of intracellular ATP was 60% on day 7. However, we observed a loss of red blood cells and RBC mass after the thawing procedure of 33% and 40% respectively.

Conclusion: With ACP 215® glycerolized and deglycerolized PRBCs meet the quality requirements for transfusion of the European guidelines, but due to the loss of RBC transfusion of 3 PRBCs is necessary to achieve the effect of 2 conventionally stored PRBCs.
P 11 Erythrocyte Loss in Case of Hemapheresis Donors: Are the Legal Guidelines Concerning Donor Protection Kept?

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Institut für Transfusionsmedizin, Universitätsklinikum Jena, Germany

Introduction: The guidelines of the BÄK (Bundesärztekammer) concerning the collection of blood and blood components specify the maximum allowed annual RBC donation volume for women up to 1,000 ml and for men up to 1,500 ml. The total donation volume cannot exceed 25 L plasma per year. An interval of at least 8 weeks must be kept between 2 whole blood donations. However, apheresis donors are allowed to donate within much shorter time intervals. The actual apheresis technique permits the traceability of blood component loss. This follow up is so far only possible with VISTA™. This software allows the management of the complete donation process, including donor selection and preparation, donation monitoring and report generation.

Material and Methods: Procedure registration for 40 donors was handled with VISTA™. Donation volumes combined with the RBC loss can typically be calculated as follows:

- Draw of blood sample for the legally required control testing: approximately 18 ml RBC/27 ml plasma.
- Residual volume of the tubing set with blood-return procedure: approximately 30 ml RBC/35 ml plasma or without blood-return procedure: approximately 95 ml RBC/112 ml plasma.

Volume of blood products:
Calculation of RBC loss was carried out with a fictitious hematocrit of approximately 95 ml RBC / 112 ml plasma.

Results:

Table 1. Volume loss female donors (n = 10)

<table>
<thead>
<tr>
<th>No donations per year</th>
<th>Plasma loss (L)</th>
<th>RBC loss (ml)</th>
<th>No incomplete proced./RBC/loss (ml)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mean: 19.5</td>
<td>8.595</td>
<td>339.6</td>
<td>1/95</td>
</tr>
<tr>
<td>Min: 17</td>
<td>5.696</td>
<td>765</td>
<td>0/0</td>
</tr>
<tr>
<td>Max: 22</td>
<td>9.590</td>
<td>1,178</td>
<td>4/380</td>
</tr>
</tbody>
</table>

Table 2. Volume loss male donors (n = 30)

<table>
<thead>
<tr>
<th>No donations per year</th>
<th>Plasma loss (L)</th>
<th>RBC loss (ml)</th>
<th>No incomplete proced./RBC/loss (ml)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mean: 15</td>
<td>10.435</td>
<td>1,151.8</td>
<td>0.65/61.75</td>
</tr>
<tr>
<td>Min: 27</td>
<td>6.664</td>
<td>1,010</td>
<td>0/0</td>
</tr>
<tr>
<td>Max: 23.2</td>
<td>13.089</td>
<td>1,592</td>
<td>3/285</td>
</tr>
</tbody>
</table>

Conclusion: With VISTA™ it is possible for the first time, to obtain watertight computer supported documentation of the donation procedures, including the blood loss volumes that occurred. The results obtained indicate that, in case of regular donations, 30% of the female and 12% of the male donors lose more RBC volume than permitted by law. Supplementary investigations are necessary to ascertain the safety of the donors in relation to the RBC loss is questioned.

P 13 Double Dose Plateletpheresis Increases Neutrophil Activation and Platelet-Neutrophil Complex Formation in Volunteer Donors

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Akdeniz University School of Medicine, Department of Hematology and Blood Bank, Antalya, Turkey

Objectives: Platelet-neutrophil complexes (PNC) might play an important role in thrombotic and inflammatory diseases. It has been shown that during extracorporeal circulation, such as haemodialysis and cardiopulmonary bypass, platelets may form heterotypic aggregates with leukocytes via platelet CD62p and leucocyte β2 integrins. There were conflicting results and limited data on the impact of plateletpheresis procedures on PNC formation and neutrophil activation on donors. In recent years, it has been possible to collect double dose platelets by new generation devices and there were no studies concerning the neutrophil activation and PNC formation on donors during double platelepheresis (DP).

Methods: In this study, we investigated the effects of DP with two different devices (Fresenius AS 204 n = 10 and MCS Plus n = 22) on in vivo neutrophil activation and PNC formation in 22 volunteer donors. Peripheral blood samples were taken immediately before and after apheresis and on the 1th, 7th days. Changes in PNC formation and neutrophils activation was determined by quantitating the CD42b+ neutrophil counts and the amount of mean flurosans intensity (MFI) of CD62L, CD54, CD50, CD11b/18, CD42b expressions by using a whole blood method on flow cytometry.

Results: Statistically significant increases were found on 42b+ neutrophil (PNC formation) percentage and counts after apheresis with both machines. CD11b/18, CD50 and CD54 expressions (MFI) on neutrophils did not show any changes after apheresis with both devices. As a marker of neutrophil activation, CD62L expression (MFI) decreased significantly after apheresis with Fresenius machine on the first and seventh days but this was not seen with MCS+ Plus.

Conclusion: Our results show that DP with Fresenius AS.TEC 204 and Haemonetics MCS+ devices results an increase on PNC formation which may be a risk factor for thrombosis and inflammation. However, clinical significance of these findings during apheresis procedures was still not known exactly.

P 18 Successful Mobilisation and Collection of Peripheral Blood Stem Cells (PBSC) after Priming with Chemotherapy and Pegfilgrastim

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Objectives: A successful collection of peripheral CD34+ cells in a 52-year-old patient with mantle cell lymphoma stage II B after priming with chemotherapy and Pegfilgrastim was performed.

Methods: The results were compared to a group of twelve newly diagnosed patients with non Hodgkin lymphoma who got daily filgrastim after chemotherapy (7×2–4 g Cyclophosphamid alone, 5× various combination treatments) for priming.

Results: Collection was performed on day 11 after the fourth treatment cycle of CHOP/Mabthera followed by Pegfilgrastim (day 2). At this time WBC count was 5.3 × 10^9/ml and 0.74% CD34+ cells in peripheral blood were detected. After one large volume apheresis (14,400 ml) 14.84 × 10^8 nucleated cells with 2.33% CD34+ cells in theuffy have been collected. In the control group the median WBC count was 10.2 × 10^9/ml, the CD34+ cells were 0.31% in peripheral blood and the nucleated cells in theuffy 10.9 × 10^9 with 1.56% CD34+ cells after a median apheresis volume of 9,000 ml.

Conclusion: Pegfilgrastim was well tolerated and save in our patient and was efficient in combination with chemotherapy as priming therapy for the mobilisation of peripheral CD34+ cells in comparison to the control group.

P 20 Impact of CD34+ Cell Dose on Engraftment in Allogeneic Non-Myeloablative PBSC Transplantations

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Masaryk University Hospital Brno, Czech Republic

Objectives: The number of cells expressing CD34+ and the number of CFU-GM can be used to assess the peripheral blood progenitor cell (PBPC) graft quality and predict engraftment in autologous and allogeneic PBSC transplantation. However, little is known about relationship between the number of transplanted hematopoietic progenitors and time to engraftment after allogeneic non-myeloablative transplantation.

Patients and Methods: Patients (18 males, 12 females) with median age of 53 years (range 27–65) were treated for AML (n = 10), CML (n = 9), lymphoma (n = 5), myeloma (n = 2), CLL (n = 2), MDS, and aplastic anemia (n = 1). Transplantations using non-myeloablative chemotherapy (busulphan, fludarabine, ATG) and sibling HLA matched PBSC were performed between 3/1998 and 12/2002. The relation between progenitor dose and engraftment (day of WBC > 1 × 10^9/l, granulocyte > 0.5 × 10^9/l, ...
and PLT > 50 x 10⁹/L) was analyzed with Spearman-rank correlation matrix. Results: A significant correlation between number of transplanted CFU-GM (median 105.8 x 10⁹/kg, range 16.0–347.7) and leukocyte as well as granulocyte engraftment was not found. On the other hand, the CD34+ cell dose (median 7.2 x 10⁹/kg, range 2.7–15.4) strongly correlated with leukocyte (r = 0.64, p < 0.001) and granulocyte (r = 0.66, p < 0.001) engraftment. Correlation was found for platelet engraftment (r = 0.55, p < 0.05) with the number of transplanted CFU-GM as well as with the CD34+ cell dose (r = -0.70, p < 0.001).

Conclusion: We can conclude, that the dose of CD34+ cells significantly affected leukocyte, granulocyte, and platelet engraftment after non-myeloablative allogeneic transplantation. However, the number of CFU-GM correlated only with platelet recovery and seems to not be useful for engraftment prediction contrary to the number of CD34+ cell, which could be used for prediction of time to hematopoietic recovery.

P 28
Red Cell Exchange in Methemoglobinemia Caused by Eugenol (Clove Oil) Intoxication – Case Report

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1Blood Center, 2Clinic of Anesthesiology and Resuscitation, University Hospital Ostrava, Czech Republic

Background: A case of 20-year-old man intoxicated by parenteral (partially paravenous) application of eugenol (clove oil) in the cubital vein is described. This chemical causes methemoglobinemia, hemolysis and its action in the organism is described as mitochondrial poisoning. Clove oil is used in dentistry as an analgesic, anesthetic and antiseptic, it is also used in perfumery. Methemoglobin is the oxidized form of hemoglobin in which the iron in the heme component has been oxidized from the ferrous to the ferric state. This renders the hemoglobin molecule incapable of effectively transporting and releasing oxygen to the tissues. The episodes of methemoglobinemia and hemolytic anaemia have been due to exposure to aromatic nitro and amino compounds (phenol, naphtalen, antimalaric agents etc).

Case Report: The patient was admitted to the hospital with consciousness alteration, cyanosis, dyspnea and high level of hemolysis and myolysis products. Deep necrosis developed in the application site in the cubita. The methemoglobin level was 17% and oxygen saturation was of 66–77%.

Controlled ventilation after intubation was started. 8 blood units of packed red blood cells were applied and then therapeutic plasma exchange was performed to eliminate myolysis and hemolysis products. Because of remaining low oxygen saturation in spite of intensive oxygenotherapy and repeated erythrocyte transfusion, it was decided to perform red cell exchange (RCE). This procedure has run from 20.00 till 23.30 o’clock. RCE is able to improve or normalize the oxygen tissue supply by removing erythrocytes containing pathological hemoglobin and supplementing functional hemoglobin for oxygen transport of the tissues. The COBE-Spectra blood cell separator was used together with Spectrastherm blood warmer. Connection of patient to the extracorporeal circulation was performed via vsuvelavia by double-lumen dialysis catheter. Anticoagulation was ensured using ACD-A solution in the ratio blood:ACD-A 15:1. Before apheresis 250 mg of Solu-Medrol i.v. were applied. The possibility of citrate toxicity was eliminated by continuous application of calcium gluconate during the whole procedure. For substitution twelve transfusion units (3,400 ml) of delecetocited and by ionizing radiation (32Gy) irradiated packed red cells were used. Red cell units no older than 7 days were used. Only 30 min after RCE has started, the first increase of oxygen saturation was noticed. During the whole procedure the hemoglobin oxygen saturation has increased continuously up to 90%. In the next few days the oxygen saturation was in the normal range.

Discussion: Although it is as yet unproven, the efficient and rapid removal of unwanted red blood cells by apheresis might also be helpful in treating rare conditions, whose pathophysiology suggests, that benefit might ensue. In the literature there are described conditions, where for instance using of red cell exchange is used in the treatment for instance malaria, babesiosis and carbon monoxide poisoning. Red cell exchange in above mentioned patient was performed in the situation, when all standard therapeutic means were used (including methylene blue application) without any success. All potentially adverse and undesirable effects associated with the massive blood transfusion were considered carefully before treatment. The procedure was accomplished without any problems and complications.

P 31
Therapeutic Approaches in the Management of Oral Cyclosporin A Intoxication

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1Department of Blood Group Serology and Transfusion Medicine, 2Department of Cardiothoracic Surgery," 3Department of Clinical Pharmacology, University Hospital, Vienna, Austria

Background: A 68-year-old male patient who was renal transplanted because of bilateral end stage kidney disease of unknown origin received a 100-fold oral overdose of CSA. Renal parameters were within the normal range. Although conventional detoxification therapy was started immediately plasma levels of Cyclosporin A (CsA) exceeded 1,500 ng/L. Because this patient was at high risk to develop renal failure whole blood exchange (WBE) as additional detoxification management was applied.

Material and Methods: For WBE a portable cell separator (MCS 3p/Haemonetics®) was used. The exchange medium consisted of 4 irradiated packed red blood cells (PRBCs) and 4 solvent detergent inactivated pooled plasmas (Octaplas®). Because of acute renal failure a continuous hemofiltration with a filtration rate of 2,000 mL/h was applied 2 h after the whole blood exchange.

Results: WBE had no immediate effect on serum CsA levels. 1,110 ng/mL were detected after the exchange procedure. 1,022 ng/mL were found in the depleted RBC fraction and 1,237 ng/mL in the depleted plasma fraction of the patient. A rapid decrease of CsA in plasma, however, was seen after starting the hemofiltration. A CsA level of 159 ng/mL was found after 72 h. The patient recovered and was dismissed from the hospital 1 month later.

Conclusion: WBE has no immediate therapeutic effect in CsA intoxication. The decrease of CsA plasma levels after onset of hemofiltration would at first sight suggest therapeutic benefit but when compared to the mean elimination half-life of CsA its effect is doubtful.

P 39
Kidney Transplant Rejection Reaction and Plasmapheresis

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Purpose: The belief that circulating antibody might be involved in rejection led to extensive use of plasmapheresis therapy. In hyperacute rejection of allografts, where therapeutic apheresis would have the firmest theoretical footing, the results wherein uniformly negative. It is apparently not possible to remove sufficient antibody quickly enough to save such grafts.

Methods: Between 1994–2002, 22 patients who exhibited symptoms of gradual rejection of kidney transplants were admitted to the apheresis treatment department of Vilnius University Hospital Santariskiu Clinic. High circulating immunological complex (CIC) levels were detected in their blood. We used the interrupted centrifugal method. We performed 6 to 7 procedures per patient with ‘Baxter corp.’ plastic bags. We removed about 3 l of plasma on average. We compensated for the lost quantity of plasma by Sol. Natrii chloridi 0.9%.

Results: See table 1, next page.

Conclusions: Plasmapheresis can be applied as supplemental method of treatment for patients following kidney transplantation in cases of kidney transplant rejection reaction. This will allow reduction of immunosuppressant doses and CIC levels in the blood.

P 41
Plasma Exchange Therapy in Refractory Myasthenia Gravis (MG) and Guillain Barre (GB) Patients

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Objective: Myasthenia gravis (MG) is a disorder of neuromuscular transmission characterized by weakness and fatigue. Guillain Barre (GB) is a common polyneuropathy causing disability and respiratory failure. Both syndromes are caused by autoantibodies directed against the postsynaptic nicotinic acetylcholine receptor in neuromuscular junction and against nerve constituents respectively. Plasma exchange (PE) in association with additional immunosuppressive agents has been employed in refractory MG/GB patients.
Severe Sharp Syndrome with Pulmonary Manifestation: Experiences with Immunoadsorption

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Introduction: A now 12-year-old patient (1.40 m, 33 kg) with Sharp syndrome, Raynaud's phenomenon, existing arterial and pulmonary hypertension (65/48 mm Hg), restrictive ventilation disorder (FEV1: 66%, MEF50: 303), restricted ventilation disorder (FEV1: 40.4%, MEF50: 74%), and pulmonary hypertension (144/104 mm Hg) was separated by Cobe Spectra peripheral venous (blood flow: 20–25 ml/min, ACD-AWB = 1:20). Immunosorba was floated continuously with 15 ml/min. A bolus of heparin (1,000 I.E.) was given into the extracorporeal system at the beginning of plasmapheresis, a permanent infusion of heparin was not necessary.

Results: ANA and autoantibody (aab) against dsDNA were reduced significantly (ANA: 1:640 to 1:160 and dsDNA-aab: 130.1 IU/ml to 42.3 IU/ml). ScI-70-aab was eluated most of all. The blood pressure was regulated (144/104 mm Hg to 105/65 mm Hg after treatment) and made a reduction of drugs possible. There is no detection of a pulmonary hypertension now, a normal heart function demonstrated by echocardiography. Lasix initial 30 mg was stopped. An improved pulmonary function is recognized (FEV1: 72.5%, MEF50: 84%).

Conclusion: The significant improvement of quality of life is in addition to the therapy with Bosentan (endothelin receptor antagonist) a success of plasmapheresis.

P 48
Therapeutic Plasma Exchange in Patients with Thrombotic Thrombocytopenic Purpura

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Introduction: Thrombotic Thrombocytopenic Purpura (TTP), first described in 1924, is an increasingly common, potentially fatal, multisystem disease process clinically defined by severe thrombocytopenia, microangiopathic hemolytic anemia, neurological abnormalities, renal dysfunction and fever. Combination of the Therapeutic plasma exchange (TPE) and plasma infusion is considered as the most suitable therapy of the disease. TPE helps to remove ULv-WF multimers and IgG inhibitor, and the addition of plasma supports the patients with the vWF-cleaving protease activity.

Methods: Authors refer their experience with 303 TPE procedures performed in 1999–2003, in 6 patients with TTP who developed heavy neurological symptoms and prolonged thrombocytopenia. The course of the disease was unexpectedly complicated with repeated early relapses. TPE procedures were performed daily by the apheresis technique Cobe Spectra in F 5 / M 1, median age 36 (19–58) years. Median volume of exchanged plasma was 0.8 (0.4–1.4) of the patient’s plasma volume which corresponded with 2,740 (900–4,600) ml. FFP and cryoprecipitate plasma were used as the replacement therapy.

Results: The course of the disease was in all 6 patients severe. After a short temporary period of improvement it was complicated with relapses with the significant decrease in the number of platelets, and elevation of LDH levels. Subsequently, it was necessary to continue in TPE procedures and to combine TPE with vincristine and cyclophosphamide. All the patients have recovered and they have no symptoms of the disease until now. The incidence of the adverse reactions like hypocalcemia, allergic reactions, tachycardia and fever were observed in 32 procedures (11%).

Conclusion: The exact cause of TTP has not yet been explained but identification of vWF-cleaving protease has led to a new understanding of the pathogenesis of TTP, and to an understanding the mechanism of the therapy using TPE. TPE is the efficient procedure that helps to patients to overcome the disease. No serious adverse reactions in patients have been observed.
Material and Methods: We evaluated 14 TTP patients, 9 females and 5 males with mean age 55 years (range 26–80 years), treated with PEX in our hospital during the last 7 years. According to the treatment protocol a mean 38 ± 5 ml plasma/kg of body weight was exchanged daily during serial logical evidence of remission appeared, defined as increase of platelet >150,000/mm³ and normal levels of LDH which was defined as response to therapy. The PEX was performed with the machine MCS+ (Haemonetics) and the plasma was exchanged for fresh frozen plasma. To assess the early response to treatment, the decline of LDH and platelet level from the first to the third cycle was calculated as: LDH before the third PEX/platelets before the first PEX (LDH ratio) and platelets before the third PEX/platelets before the first PEX (platelet ratio). Statistical analysis was performed with Wilcoxon test.

Results: We found that 71.5% of the patients responded to therapy with PEX. Four patients were non responders (28.5%). After 2 procedures of PEX none clinical or laboratory marker correlated with outcome. After 2 procedures of PEX only LDH and platelet level had improved in responding patients. The LDH ratio was the best predictive marker of response (p<0.01). Nine of 10 responding patients had an LDH ratio <0.60 and all of the non responders patients had LDH ratio > 0.60. Conclusion: TTP remains a serious condition and the early identification of patients not responding to PEX might help to modify the therapy with intensive plasma exchange and additional therapeutic interventions. The LDH ratio might be a useful predictive marker for separating responding from non responding patients to plasma exchange.

P 52
Thrombotic Thrombocytopenic Purpura (TTP) and Thrombophilia – Congenital or Acquired?

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Purpose: We investigated our TTP patients for the existence of thrombophilic disorders congenital or acquired contributing to more aggravate and relapsing forms of TTP.

Material and Methods: The last 7 years a total of 21 patients 8 male and 13 female with mean age 46 years and the established diagnosis of TTP were treated in our hospital. Fourteen patients had the relapsing form of the TTP. All the patients entered the protocol with methyl prednisolone and plasmapheresis exchange (PE) by using the machine MCS+ (HEAMONETICS). Twelve out of 21 patients were studied for thrombophilic disorders: FXIII activity, Protein C, S, ATIII, APC-Resistance (APC-R), genotyping analysis for FVLeden/FI20210/MTHFR, plasminogen, Anticardiolipin antibodies (ACA) and Lupus anticoagulant (LA). The statistical analysis was performed with χ² test.

Results: We found 7 patients with positive markers of thrombophilia. The characteristics of the 7 patients are shown in the table.

<table>
<thead>
<tr>
<th>Patient</th>
<th>Sex</th>
<th>Age</th>
<th>Relax</th>
<th>Number of PE</th>
<th>Coexisting disease</th>
<th>Thrombophilic disorder</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>M</td>
<td>44</td>
<td>yes</td>
<td>19</td>
<td>infection of the skin</td>
<td>FVLeden</td>
</tr>
<tr>
<td>2</td>
<td>F</td>
<td>63</td>
<td>no</td>
<td>16</td>
<td>cancer of the bladder</td>
<td>LA +</td>
</tr>
<tr>
<td>3</td>
<td>M</td>
<td>58</td>
<td>no</td>
<td>8</td>
<td>–</td>
<td>PrS↓</td>
</tr>
<tr>
<td>4</td>
<td>F</td>
<td>71</td>
<td>no</td>
<td>18</td>
<td>–</td>
<td>MTHFR</td>
</tr>
<tr>
<td>5</td>
<td>F</td>
<td>50</td>
<td>no</td>
<td>13</td>
<td>autoimmune</td>
<td>ACA IgM+</td>
</tr>
<tr>
<td>6</td>
<td>F</td>
<td>59</td>
<td>yes</td>
<td>7</td>
<td>–</td>
<td>FVLeden</td>
</tr>
<tr>
<td>7</td>
<td>F</td>
<td>22</td>
<td>no</td>
<td>8</td>
<td>–</td>
<td>FVLeden</td>
</tr>
</tbody>
</table>

M = Male; F = female.

The patients 1M with FVLeden and 4F with low levels of PrS and MTHFR needed >15 procedures in order to recover completely. The patient 3M with low levels of PrS and 6F with FVLeden presented DVT (Deep Vein Thrombosis) one week after the last PE. The two patients 2(F), 5(F) had LA and ACA IgM positive and presented thrombosis and infection of their catheter. The patient 7F was homozygote FVLeden, he presented stroke and is on oral anticoagulant therapy without TTP relapse. The other 5 patients with negative thrombophilic tests presented with no signs of thrombosis and milder TTP presentation (p<0.001).

Conclusion: TTP is a syndrome affecting predominantly females (13/8 F:M). The confirmation of thrombophilic disorders may be crucial in therapeutic management due to the possibility of thrombosis and more aggressive presentation. The patients with positive markers of thrombophilia should be monitored closely and anticoagulant or antiplatelet therapy should be administered in cases of thromboembolic events.

P 53
Thrombotic Thrombocytopenic Purpura and Pregnancy

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Objectives: A series of women with thrombotic thrombocytopenic purpura (TTP) associated with pregnancy is presented. The study analyses the relationships between TTP and pregnancy and maternal and fetal outcomes.

Patients and methods: 40 consecutive patients who met the classic criteria for TTP were studied. Among them, 8 patients who were either pregnant or in the postpartum period were identified (20%). All patients were treated with prompt plasma exchange. A number of adjunt therapies also were used in patients not responding to plasma exchange. These included corticosteroids, antiplatelet aggregation drugs, vincristine and splenectomy.

Results: The study population consisted of 9 women, 6 of whom presented an acute single episode associated with pregnancy and 2 having a chronic relapsing form of the disease. These two patients had a total of 18 TTP episodes, 2 of which were related to pregnancy. None of them were diagnosed during pregnancy or in the postpartum period. Another pregnant patient had recurrent disease and neither episode was associated with pregnancy. Five women were pregnant and 3 were in the immediate post-partum period. In 2 patients the disease developed before midpregnancy. One gestation was successfully terminated with a normal fetus at the 26th week because of worsening of the TTP and the other by elective abortion at the 16th week of gestation. In 3 patients, the disease manifested in the third trimester and in one of these the gestation was terminated after demonstration of the death of the fetus. The other two mothers gave birth to preterm infants. Fetal mortality was 40% in pregnancies associated with TTP. There was one maternal death. This mortality rate is similar to that observed in the rest of patients of our series (4/32). There were no serious long-term sequelae in the 8 survivors. Mean follow-up of the eight survivors is 6.16 years (range 1.32–11.4 years).

Conclusion: This limited experience suggests that pregnancy could be one of the initiating agents of TTP. However, in those patients where PTT was first diagnosed during the course of a pregnancy, no further episodes were observed. Pregnancy in patients with a previous history of recurrent TTP may increase the risk for recurrence, but it can also be uncompli-

As far as survival, pregnancy does not increase the risk for the mother with TTP, when compared with the mortality rate in the rest of patients in our series. Preterm delivery and intrauterine fetal death were frequent complications of these pregnancies.

P 54
Patient with Relapsing Severe Thrombotic Thrombocytopenic Purpura and Systematic Lupus Erythematosus – A Case Report

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Objectives: Thrombotic thrombocytopenic purpura (TTP) is a devastating microvascular occlusive disorder associated with an acquired or congenital deficiency of von Willebrand factor – cleaving metalloproteinase (ADAMTS13) which normally prevents the entrance of unusually large VWF multimers into plasma. The acquired form of TTP is caused by autoantibodies against VWFcp, whereas homozygous or compound heterozygous
mutations of the ADAMTS13 gene are responsible for the recessively inherited TTP.

Case Report: The authors report the case of a 35-year-old female with systemic lupus erythematosus (SLE) and acquired form of relapsing TTP. SLE with a biopsy verified lupus nephritis IV was diagnosed 5 years before onset of TTP. The patient was treated with prednisolone, cyclophosphamide, azathioprine, and, by last year, also with cyclosporine A (initiated for the nephrotic syndrome > 10 g per day). During a regular out-patient examination only several haematomas were observed and the patient complained of fatigue. A week later, moderate anaemia, severe thrombocytopenia, scistocytes, and high level of LDH were found out, but she rejected hospital admission. Next day, neurological abnormalities were observed and treatment with fresh frozen plasma was immediately initiated. But, after all, respiratory failure, heart failure, and a systemic shock developed in few hours. A large volume plasma exchange (TPE) was promptly started even the patient was in a critical condition. TPE followed for 10 days with rapid improvement of patient’s condition and laboratory parameters. Two weeks after dismissing from a hospital, relapse of thrombocytopenia and haemolytic anaemia occurred. However, TPE was not satisfactory and splenectomy was performed resulting in a recovery of TTP.

Conclusion: The acquired form of TTP in SLE patients can be a rare but very serious complication with difficult differential diagnosis, which includes catastrophic antiphospholipid syndrome, disseminated intravascular coagulopathy, haemolytic-uremic syndrome, and activity of SLE. In respect of a frequently very fast and serious development of this disease, early initiation of the large volume plasma exchange can be a life saving therapeutic approach in these patients.

P 55 Extracorporeal Photopheresis in Patients with Chronic GVHD

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Introduction: Chronic GVHD is considered as the most common late complication of allogeneic hematopoietic stem cell transplantation. It continues to be a significant cause of morbidity and mortality. Patients are treated by the administration of conventional immunosuppressive therapy over prolonged periods of time. Nevertheless, the therapy causes significant side effects with increased morbidity and mortality due to the infections. The situation may be complicated especially in patients who are refractory to immunosuppressive treatment. New ways of therapy are searched, and promising therapeutic strategies include extracorporeal photopheresis (ECP).

Methods: We tried to evaluate the clinical effect of 167 ECP procedures in 6 patients with chronic GVHD. We assessed also the influence of the ECP on levels of T cell subsets (CD3+, CD4+, CD8+, IRI), B cells and NK in peripheral blood and in MNC concentrates. The ECP procedures were performed by means of Cobe Spectra, either in semi-automated or automated mode (v. 5.1.6). We processed 2 (1.6–2.1) total blood volumes of the patients, which corresponded with 7.1 (6–7.6) liters of blood. Photo-modification of MNC was performed by the use of 8-MOP (Gerot) and the UV-A irradiator (Psorilux 3070, Heraeus). The UV-A irradiator (Psorilux 3070, Heraeus) was promptly started even the patient was in a critical condition. TPE followed for 10 days with rapid improvement of patient’s condition and laboratory parameters. Two weeks after dismissing from a hospital, relapse of thrombocytopenia and haemolytic anaemia occurred. However, TPE was not satisfactory and splenectomy was performed resulting in a recovery of TTP.

Conclusion: The acquired form of TTP in SLE patients can be a rare but very serious complication with difficult differential diagnosis, which includes catastrophic antiphospholipid syndrome, disseminated intravascular coagulopathy, haemolytic-uremic syndrome, and activity of SLE. In respect of a frequently very fast and serious development of this disease, early initiation of the large volume plasma exchange can be a life saving therapeutic approach in these patients.

P 56 Treatment of Chronic Graft versus Host Disease of the Lung with Extracorporeal Photopheresis – Experience in 2 Patients

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Purpose: Photopheresis (extracorporeal photochemotherapy, ECP) appears to be an effective treatment for skin and liver manifestations of chronic graft versus host disease (cGVHD) in many studies. Experience in the treatment of pulmonary GvHD with ECP is still limited and controversial.

Methods: We report on 2 patients with severe pulmonary GVHD who were treated with ECP using the UVA-XTS-device (Therakos Company, Exton). Patient 1 was a 57-year-old woman, diagnosed with AML M4 8/98. Patient 2 was a 39-year-old man with AML M2, diagnosed 9/1999. Both patients were transplanted from HLA-identical siblings and suffered from severe lung and moderate liver GvHD. ECP was started 12 and 17 months after transplantation, respectively. Additional immunosuppressive treatment consisted of cyclosporine, mycophenolat mofetil and prednisolone (30mg/kg) in Pat. 1 and tacrolimus and prednisolone (30mg) in Pat. 2. ECP treatments were performed on 2 consecutive days every week during the 1st month and every 2nd week thereafter for the following 5 months.

Results: Treatment intervals were tapered thereafter individually according to patients condition.

Conclusion: The ECP procedures were well tolerated by both patients without any severe side effects. Technical problems were mostly related to difficult venous access. Patient 1 was treated with ECP for 1 year, during which time pulmonary symptoms improved continuously and liver function parameters normalized. Now she is still in stable clinical condition 2.5 years after completion of ECP treatment. Patient 2 has been treated with ECP for 20 months and therapy is still ongoing. His respiratory symptoms and liver function tests improved substantially. In both patients immunosuppressive medication could be reduced and steroid dosage could be tapered to 5mg. Symptoms of Cushing disease completely resolved.

P 57 Extracorporeal Photoimmunotherapy (ECP) for Treatment of Steroid Refractory Extensive Chronic Graft versus Host Disease (cGVHD): Turkish Experience

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Objectives: cGVHD can arise as a late complication after allogeneic hematopoietic stem cell transplantation. Patients with extensive disease to date require intensive early and long-term immunosuppression. We aimed to share the experience of three Turkish hemapheresis centers using ECP in this complicated patient population.

Methods: Thirteen patients in three centers were treated with ECP (UVAR-XTS) on 2 consecutive days every 2–4 weeks until resolution of GvHD over a period of 6–15 months. Beyond extensive cutaneous cGVHD, four patients had also bronchiolitis obliterans (BO). Skin scores assessed by an experienced dermatologist. Laboratory and radiological findings after 4 months of ECP were accepted as response criteria. In this almost fully automated system mean 261.4 mluffy-coat was processed within 193 min using UVADEX sterile solution.

Results: After a median of 12 cycles of treatment, all patients showed a favorable response. ECP was tolerated well only one patient developed Gr4 thrombocytopenia and another patient had a massive GIS bleeding due to...
an esophageal tear. Reduction in cholestatic parameters was observed in 9, improvement in respiratory functions and CT evaluations in 4, and reduction in immunosuppressive agent need in 10 patients. The most impressive result was the reduced need for hospitalization of these patients and improvement of skin lesions. All but one of the skin biopsy scores was also better after ECP. Conclusion: As chronic extensive GvHD is a life devastating disorder, every attempt to improve the quality of life should be evaluated carefully. Our findings suggest that ECP is a safe and effective adjutative therapy for steroid refractory chronic extensive GvHD of the skin. The place of ECP for the treatment of visceral GvHD and additional co-morbid conditions like BO should be further analyzed in large scale trials.

P 65
Hemotherapy in Kidney Transplantation in the Military Medical Academy
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Objectives: A total of 107 kidney transplantations (81 males and 26 females) from living donors (group I) and 14 kidney transplantation (8 males and 6 females) from cadavers (group II) were performed in the period 1996–2002 at the Military Medical Academy (MMA) in Belgrade in the Serbia and Montenegro.

Methods: Lymphophytotoxic cross-matching was done before each kidney transplantation. In perioperative transfusion treatment determined quantity of filtered red blood cells (F-RBCs) and/or filtered platelets (F-PLT) were given to recipients according to blood loss.

Results: In only 9 (8%) recipients in group I transfusion therapy was not applied perioperatively. An average use of 2.02 units of F-RBCs (942.70 ml) was used intraoperatively in 67 (63%) recipients in group I, an average of 1.9 units of F-RBCs (521 ml) was used before kidney transplantation in 15 (14%) recipients in group I and an average use of 4.06 units of F-RBCs (1 208.68 ml) and an average of 0.63 unit of F-PLT were used postoperatively in 61 (57%) recipients. In all recipients from group II used transfusion therapy was applied perioperatively. An average use of 2.02 units of F-RBCs (942.70 ml) and an average of 1.4 units of F-PLT were used postoperatively.

Conclusion: All recipients well tolerated the therapy and no adverse effects of the therapy were observed. The need for transfusion therapy intraoperatively was approximately same in both recipient groups, while in recipients from cadavers need for transfusion support in posttransplantation period was much higher.

P 68
Optimization of Therapeutic Procedure During LDL Apheresis – a Multivariate Computerized Model
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Objectives: LDL apheresis is a very effective method in the treatment of resistant hypercholesterolemia when other therapy (dietary, medicamentous) fails. This is essential and life saving in homozygous hypercholesterolemia, in other cases it might substantially prolong the life without premature occurrence of severe atherosclerotic complications. To maximize efficacy of the usage of LDL absorbers we aimed to create a multivariate computerized model.

Methods: LDL apheresis was performed (after separating plasma in a continuous-flow blood separator Cobe Spectra, USA) by absorption-desorption automat ADA (Medicap, Germany) controlling the passage of plasma through a pair of columns containing Sepharose 4b carrying an anti-apoprotein B antibody (Lipopak, Focard). Computerized model was created to control the volume and flow of plasma for optimal adsorption performance. Based on the pilot studies the following data were used for calculation: height, body weight, sex, baseline and expected plasma LDL cholesterol. The principle of calculation is an exponential dependence and suggestion that the absorber eliminates all LDL cholesterol and that the amount of eliminated LDL cholesterol correlates with the binding column capacity. Presumption is even distribution of LDL-cholesterol in plasma, total elimination during passage through column, permanent steady of absorption column and zero neglected release of LDL cholesterol into circulation from tissue reserves during the 4 h of procedure.

Results: Suitable software has been created, using Microsoft Excel, including HTML version. User loads the above mentioned data (see formula) and based on the results of calculation changes optimal plasma flow through separate columns to gain the expected goal. On the contrary to the previously used empiric method a shortening of the procedure has been reached, which has both positive medical and financial consequences.

Conclusion: Software for calculation based on the above presumption has been created and validated in the first twenty procedures. It enables to rationalize setting of LDL apheresis, with maximal efficacy of absorption columns. Recently, the software is being tested in a larger number of procedures.

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