were evaluated and compared with the theoretical energy requirements.

**Results** Mean caloric intake on the first day was 59% (SD±46.4) and on the second day was 64.3% (SD±47.4) of daily requirements. 72(57.1%) patients received ≥80% of required daily calories, without a significant difference with patients who received <80% of daily calories, (p=0.3).

The overall mortality rate was 28.5%. Patients who received <80% of daily caloric needs were 4 times more likely to have a fatal outcome, compared to those who received ≥80% of daily caloric needs [OR=4.0 95% CI (1.2–12.7) p=0.01].

Daily caloric intake of ≥80% resulted a protective factor against death in the Cox proportional-hazard regression model (β= –1.1, p=0.02).

**Conclusions** We have to increase the number of patients who receive ≥80% of daily caloric requirements and provide appropriate nutritional support during the first days of admission. Mortality rate remains high, due to the large number of patients receiving <80% of needed calories.

**1464 MEDICAL RADIATION EXPOSURE IN CHILDREN DIAGNOSED WITH ACUTE LYMPHOBLASTIC LEUKEMIA FROM 1995–2010: A SINGLE INSTITUTION STUDY**

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**Objective** This retrospective study examines a cohort of children diagnosed with acute lymphoblastic leukemia, examining exposure to medical radiation pre-conception, pre-natal or in early childhood. Exposure is documented through family interview. The study encompasses children diagnosed with A.L.L. and treated at the Children’s Hospital of Pittsburgh over a fifteen year period.

**Background** Early exposure to medical radiation is one of the identified risks for childhood leukemias but documentation is difficult and mostly lacking in the United States experience. The author of this study developed a questionnaire that examines radiation exposures in either parent of to the child later diagnosed.

**Methods** Each family who was consented to be interviewed completed a five page questionnaire at clinic visit, through phone or mail. Whenever possible both parents were interviewed.

**Results** To date the author has been able to interview about 70% of children diagnosed from 2005–2010 however the interview rate for the period 1990–2005 is approximately at 5%. Among the families interviewed at least one exposure was commonly documented.

**Conclusions** Exposure to medical radiation for a child later diagnosed with A.L.L. may at occur at several critical junctures. Chest or sinus x-rays or CT of a parent pre-conception, particularly repeated scans have the possibility of DNA damage. Early childhood exposure through the diagnostic process (ruling out infection or trauma) may well contribute to this “perfect storm” in the still elusive causes of childhood A.L.L.

**1465 CLINICAL CHARACTERISTICS AND TREATMENT RESULTS OF NEUROBLASTOMA PATIENTS**

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**Objective** and method: The survival of the patients with neuroblastoma has improved in last few decades. But still it depends on various clinical and biological factors. To assess the clinical features and trends in survival, the data for 56 newly diagnosed patients between September 1996 and August 2011 from a single center were retrospectively analyzed.

**Results** Histopathologic subtypes were neuroblastoma (NBL) in 52 patients and ganglioneuroblastoma in 4 patients. The median age was 2.5 years and Male/Female ratio was 1.2/1. Primary tumor sites were abdomen, thorax, and neck with the frequency of 77.4%, 19% and 3.6% respectively. There were 21, 22, 9, 31 patients with stage 1, 2, 3, 4, 4S disease and their 5-year survival rates were 100%, 74%, 33%, 6.9%, and 59%, respectively In multivariate analysis, stage 4 disease (P<0.001), abdominal primary tumor site (P<0.001), NBL subtype in histopathology (P=0.001), and responsiveness to chemotherapy (P<0.001) were the determinants of poor prognosis.

**Conclusions** The survival rates in children with local disease are comparable with the results of developed countries; however, the results in children with advanced disease are still not satisfactory. To improve the outcome, especially in children with advanced disease, more effective chemotherapy regimens and molecular therapies should be investigated. Sharing the knowledge and capacity building to improve the treatment results in NBL are also critical for developing countries.

**1466 THE ETIOLOGY OF PERIPHERAL LYMPHADENOPATHY IN CHILDREN**

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**Aim** To determine the clinical and laboratory findings of peripheral lymphadenopathies.

**Methods** We evaluated 224 children who had been admitted to our pediatric clinic because of enlarged peripheral lymphadenopathy (LAP). The patients age, sex, localization, duration, size, mobility, extension of lymph nodes, systemic symptoms, diagnostic laboratory and radiologic investigations and final diagnosis were determined. Features of lymphadenopathy were compared between patients with malignant and benign diagnosis.

**Results** The ages of the patients ranged between 2 months and 16 years (median 7 years); 56% of the patients were male. Of the 186 (83%) patient had benign and 38 (17%) had malignant disorder. A total of 164 (73%) had localized, 60 (27%) had generalized LAP. The most frequent cause in the benign group was acute lymphadenitis where as hodgkin lymphoma was most frequent in the malignant group. Cervical region was the most common localization for benign or malign disorder. The mean age was higher in the malignant group. We determined acute LAP in 164 (73%) and chronic LAP in 60 (27%) patient. We didn’t find differences between the benign and malignant groups according to size and mobility of nodes. Anemia, leukaopenia trombocytopenia and organomegaly were significantly common in malignant group. Weight loss, night sweet and fatigue were more frequently associated symptoms in the malignant group. Excisional biopsies were performed to 50 (22.3%) patient.

**Conclusion** The following findings were important to alert the physician about the probability of a malign disorder: older age, supraclavicular lymphadenopathy, abdominal LAP, abnormal complete blood count and organomegaly.

**1467 PEDIATRIC ONCOLOGY PATIENTS PRESENTING WITH SPINAL CORD COMPRESSION**

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Spinal cord disease in children with known or suspected malignancy is an oncological emergency because it commonly implies malignant spinal cord compression. The records of 17 children with cancer presenting with spinal cord compression, encountered over
15 years, were reviewed. 11 patients were neuroblastoma, 3 were non hodgkin lymphoma and the last were rhabdomyosarcoma. Male/female ratio was 1.2, the median age was 5.2 years, 9 patients went to surgery for tumour resection and laminectomy, others were treated with chemotherapy after biopsy. After treatment, 3 of severely affected 6 patients, who presented with paralysis, completely recovered neurologic function. 2 of 4 patients with moderate deficits, consisting of paresis and bowel/bladder dysfunction, completely recovered neurologic function. 7 children, who had mild symptoms comprised of paresis alone, fully recovered.

The frequency of complete neurologic recovery in children with intraspinal mass inversely correlated with the severity of the presenting neurologic deficits. The time passed between the first symptom and treatment was also important for neurologic recovery.

**Results**

Survivin, an apoptotic inhibitor, is overexpressed in various types of cancer. It has been shown that single nucleotide polymorphisms in the survivin gene promoter can modulate survivin expression and consequently influence the risk for some cancers. The aims of the present study were to:

a. analyze by means of PCR/RFLP, genotype and allele frequencies for the promoter –31 G/C polymorphism in the survivin gene of 59 Wilms tumour (WT) patients and 82 controls,  
b. determine cytoplasmic and nuclear survivin expression in WTs using immunohistochemical methods.

The frequencies of alleles and genotypes were significantly different between patients and controls for the –31 G/C polymorphism. Individuals with CC and CG genotypes had significantly decreased risk of WT compared to GG individuals (OR 0.30, 95% CI 0.15–0.60). A statistically significant difference in cytoplasmic survivin expression between lower and higher grades risk of WT compared to GG individuals (OR 0.26, 95% CI 0.07–0.96; p=0.000), but without correlation with the genotypes. Our findings suggest that both survivin genotypes and survivin expression, though not showing direct relationship, represent relevant risk/prognostic markers for WT in Serbian population.

**Conclusions**

Incidence Overall 0.02 per 100,000 versus 2 per 100,000 worldwide.
Platelet count: Abnormally high in 57% patients.
Survival rate: Overall 83.5% (10/12) versus 70% worldwide.