We report clinical course and genotype of four patients with hepatocerebral form of MDS. Patients 1 and 2, daughters of consanguineous Roma parents, presented with liver failure at six and two days of life, respectively. The older sibling had lactic acidosis and progressive liver failure, without clear neurological involvement. Liver histology revealed giantcellular hepatitis, fibrosis, cholestasis, hemosiderosis and steatosis. Working diagnosis was neonatal hemoiserosis. Despite the treatment, disease progressed to death at 40 days. Younger sibling had similar clinical course and MDS was suspected. Immunohistochemical staining of the deceased sibling’s liver showed combined respiratory chain deficiency. Homozygous variant in the DGUOK gene in both parents confirmed the diagnosis. Despite cofactor/antioxidant treatment, patient 2 died at the age of two months.

Patient 3 presented with recurrent nonketotic hypoglycemia, cholestasis and hypotonia at the age of two months. Liver disease was slowly progressive, with permanently elevated lactate and alanine. Histology showed giantcellular hepatitis, fibrosis, cholestasis, hemosiderosis, polymorphous mitochondria and microvesicular steatosis. MDS was suspected, but immunohistochemical staining was uninformative. Due to end-stage liver disease, LTx was performed at the age of six months. Patient died in early postoperative period. Whole exome sequencing (WES) revealed biallelic mutations in the MPV17 gene.

Patient 4 had intrauterine growth retardation, severe hypotonia and developmental delay since birth. Acute liver failure, presenting with ketotic hypoglycemia, lactic acidosis, hepatomegaly and coagulopathy, occurred at the age of four months. The individual additionally developed mystagmus. Brain MRI was normal. Liver biopsy showed steatosis and abnormal mitochondria. Immunohistochemistry and clearly decreased mtDNA copy number per nuclear genome in liver pointed to MDS. The disease progressed rapidly and patient died three weeks after admission. WES revealed two biallelic mutations in the POLG gene.

Revealing genetic basis of liver failure due to MDS, with WGS as an important option, is pre-requisite to decision on LTx. It is also essential for genetic counseling and prenatal diagnosis in future pregnancies.

In order to reach goals of this study, we included 127 newborns with congenital heart disease and 103 mothers of affected newborns with congenital heart disease. We measured AdoMet and AdoHcy in their plasma by high performance liquid chromatography tandem mass spectrometry and then calculate their methylation potential and compare them with the reference values.

In the group of newborns with congenital heart disease we have found statistically significant increased AdoMet compared to the referral group (289 vs. 184 nmol/L). There was no statistically significant difference in the concentration of AdoHcy (67 vs. 81,5 nmol/L). Methylation potential in this group was consequently statistically significantly higher (4,34 vs. 2,35). In the group of mothers of newborns with congenital heart disease we have found statistically significant increase of AdoHcy compared to the referral group (20 vs. 15,2 nmol/L), but no statistically significant difference in the concentration of AdoMet (81 vs. 81,9 nmol/L). Methylation potential was statistically significantly lower (4,55 vs. 5,54).

Increased concentrations of AdoMet or AdoHcy measured in our groups of examinees and statistically significant differences in the methylation potential values in our groups of examinees compared to the reference values point to changed methylation processes which could contribute to the pathogenesis of congenital heart disease. Further studies are needed to elucidate the exact mechanism by which disturbed methylation leads to higher risk for congenital heart disease and changes in methylation processes in the fetus.

### Neonatology

**HEMMEGALENCEPHALY AND OHTAHARA SYNDROME CAUSING NEONATAL SEIZURE – A CASE REPORT**

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Hemmegalephalhy (HME) is rare congenital, hamartomatous malformation of the brain cortical development characterised by enlargement of all or a part of one cerebral hemisphere. It is estimated that prevalence range is from 1 to 3 cases per 1000 children with epilepsy, and 1-14% among children with disturbed cortical development. Ohtahara syndrome is a rare infant epilepsy syndrome, characterised by “burst supression” pattern in EEG (high amplitude spikes followed by little brain activity or flattening of the brain waves). It is accompanied by severe neurodevelopmental delay, presumably caused by HME as a structural malformation.

We present a case of female premature newborn born from bichorionic, biamniotic twin pregnancy, who was delivered vaginally after 35 weeks and 6 days, as the first twin. Apgar scores after 1 and 5 minutes were 10, physical examination was uneventful. The first in vitro fertilization resulted in twin pregnancy of 27-year healthy mother. Apart from oligohydramnios before delivery, pregnancy was uneventful. Family history of both parents was unremarkable. At the age of 24-hours baby girl developed first tonic spasms lasting one minute and spontaneously ceasing, without losing consciousness.

The low availability of folic acid or vitamin B12 in the early pregnancy and consequent mother’s hyperhomocysteinemia are associated with the risk of having a child with congenital heart disease. The goal of this study was to determine whether homocysteine itself is a risk factor for congenital heart disease or changes in methylation biomarkers S-adenosylmethionine (AdoMet) and S-adenosylhomocysteine (AdoHcy) and methylation potential are responsible for a higher incidence of congenital heart disease.

**METHYLATION BIOMARKERS S-ADENOSYLMETHIONINE AND S-ADENOSYLMETHIONINE AND METHYLATION POTENTIAL IN NEWBORNS WITH CONGENITAL HEART DISEASE AND THEIR MOTHERS**

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She received intravenous Phenobarbital 10 mg/kg followed by 5 mg/kg/day.

Initial laboratory findings including lumbar puncture and initial metabolic evaluation were all unremarkable. A tumor or congenital malformation of the left hemisphere of the brain was suspected after the first two dimensional brain ultrasonography. Multi Slice Computed Tomography (MSCT) revealed left HME, confirmed with the magnetic resonance imaging (MRI) together with polymicrogyria of frontal lobe, atypical form of the left Sylvian fissure and the left frontal ventriculomegaly. She developed refractory seizures (tonic; focal with automatisms – squelching, eye blinking; generalised, often waking her up from sleep). EEG showed suppression burst pattern and after extensive diagnostic evaluation the Ohtahara syndrome was diagnosed.

Despite several different antiepileptic drugs, and their different combination, frequency and severity of the seizures did not improve and she developed severe developmental delay. At the age of 10 months she underwent functional hemispherotomy, and so far, eight months after the surgery she experienced no seizures together with major improvement in neuromotor development (despite strabismus and right hemiparesis which occurred after surgery). Her twin sister is healthy, normally developing, without seizures. Our findings are in comply with the data from the literature, claiming that after surgery the improvement of the patients is remarkable.

Results From the opening to mid-February, 135 women showed interest in donating human milk, of which 28 became donors.

In February 2020, we had 24 still active donors. Four women stopped donating, with a median donation period of 2 months. All donors were tested for blood borne viruses by serology and NAT and were negative. In total, we received 79 L of donated human milk. We started pasteurizing the milk after obtaining a license. Of the 30 controlled pre-pasteurization milk pools, 10% was over the allowed microbial contamination. Microbiological controls were performed for each pasteurization cycle. They were all sterile. The nutritional values of milk were all within the expected range.

Conclusions Human milk is recognised as the optimal feeding for all newborn infants.

When mothers’ own milk is not available, donor human milk provided by HMB is the second-best choice, especially for premature or sick infants. The opening of the HMB in Croatia is highly important in helping to provide the best possible medical care for prematurely born babies and infants with a serious medical condition when they cannot receive their mother’s milk.