regulated by a numerous enzymes that modify and transfer sugar residues to amino acid side chains.

Over 150 different types of CDGs have been described. The most common are phosphomannomutase 2 (PMM2-CDG) and -1,3-glucosyltransferase deficiency (ALG6-CDG). Symptoms common to all CDGs are seizures, psychomotor delay, hypotonia, feeding disorders, liver disease and coagulopathy. Diagnosis is based on clinical presentation and sialotransferrin profiling and confirmed by gene analysis. Early diagnosis is critical in disorders for which specific therapy exists.

Female infant was born from a third, uneventful pregnancy. Parents are not related and older siblings are healthy. Since birth, the infant was severely hypotonic and exhibited developmental delay, feeding disorder and failure to thrive. At the age of three months she was transferred to our Department for further work-up. At that time, she had low body weight (4314 g, 1st c) and was hypotonic with reduced spontaneous movements. We had noticed dysmorphic features- hypertelorism, micrognathia, abnormal ears, flattened nose, rhizomelic limbs, inverted mamillae and abnormal fat distribution. She was fed with elemental infant formula through the nasogastric tube. The patient had small intestine malrotation that manifested with recurrent vomiting which was resolved after surgical intervention. She developed thrombosis of the brain venous sinuses due to coagulopathy (low concentration of antithrombin III, protein C and coagulation factors IX and XI). She was also diagnosed with hypothyreosis and suffered from generalised oedema with hypoalbuminemia as a consequence of protein losing enteropathy, treated with albumin infusions. Protein losing enteropathy improved on elemental formula.

Severe epilepsy was drug-resistant. Sialotransferrin profiling pointed towards a glycosylation defect (elevated di-sialotransferrins, lowered penta- and tetra-sialotransferrins). Gene analysis revealed biallelic mutations in the ALG6 gene, which codes for glycosylation enzyme -1,3-glucosyltransferase.

ALG6-CDG has a recognizable phenotype characterised by hypotonia and proximal muscle weakness, extrapyramidal signs, pharmacoresistant epilepsy, coagulopathy, protein losing enteropathy and dysmorphic features. Described patients had poor prognosis. There is no specific therapy available. Early diagnosis is important to predict and treat symptoms and to stop diagnostic odyssey. It enables prenatal diagnostics in future pregnancies.

**106 DIVERSITY OF CLINICAL PHENOTYPE OF PATIENTS WITH PYRUVATE DEHYDROGENASE DEFICIENCY DUE TO PDHA1 GENE MUTATIONS**

Katarina Škocic*, Danijela Petkovic Ramadža, Tamara Žigman, Nina Barilić, Ivan Lehman, Johannes A Mayr, Holger Proksch, Sarka B Wirtmann, Wolfgang Spel, Nikola Mesarić, Valentina Rahelić, Ksenija Fumić, David Özretić, Maja Tomasović, Ivo Barić. Department of Pediatrics, University Hospital Centre Zagreb, Zagreb, Croatia

Pyruvate dehydrogenase (PDH) is a mitochondrial multienzyme complex which catalyzes pyruvate oxidation (derived mainly from glucose) to acetyl-CoA which then enters tricarboxylic acid cycle. Its deficiency results in accumulation of lactate and alanine. Lactate/pyruvate ratio is normal. Brain is almost always affected because it obtains energy primarily from oxidation of glucose. PDH deficiency is most commonly due to mutation in an X-linked pyruvate dehydrogenase E1 alpha subunit gene (PDHA1). Diagnosis is a challenge especially in females not only because of variable clinical phenotype but also because of skewed X-chromosome inactivation - its pattern varies among individuals, but also among different tissues of the same individual. Therefore, measuring PDH activity is not a reliable guide to the diagnosis in females. Gene analysis may be used to confirm the diagnosis.

Causal treatment options include thiamine (cofactor of PDH), ketogenic diet and dichloroacetate. The goal of the study is to alert medical professionals to clinical variability of PDHC deficiency and thus make easier early diagnosis.

We present a spectrum of clinical phenotypes of six patients with PDH deficiency diagnosed and treated in our Center.

In patients 1 and 2 disease started antenatally – ultrasonography revealed brain ventriculomegaly. After birth MR demonstrated agenesis of corpus callosum, ventriculomegaly, cerebellar hypoplasia. Afterwards those patients had seizures and severe psychomotor retardation. Patient 3 had conatal hypotonia, respiratory difficulties and severe lactic acidosis. MRI revealed mild diffuse brain atrophy. At 8 years he has generalized hypotonia, walks with support. At age 6 months patient 4 presented with somnolence, metopic synostosis and mild white matter atrophy and ventriculomegaly. At 6 years she has hypotonia, moderate psychomotor retardation and ataxia. Patient 5 presented at 3 months with acute encephalopathy of unclear etiology. At 7 years she has isolated epilepsy and normal psychomotor development. Her older brother died at the age of 17 days due to cardiorespiratory failure of unknown cause. Patient 6 have had intermittent ataxia since 1 year and 8 months of age. Now at 10 years he has normal cognition. All patients have elevated plasma alanine and lactate (2,66-19,9 mmol/L; ref. range 0,63-2,40), and CSF lactate (3,64-8,52 mmol/L; ref. range 1,2-2,1), with normal lactate/pyruvate ratio. All patients were on ketogenic diet and thiamine treatment. No patient had regression or deterioration while on therapy.

PDH deficiency should be in differential diagnosis of most neurological signs and symptoms. Early diagnosis may enable optimal treatment and proper genetic counseling.

**107 EARLY ONSET LIVER FAILURE DUE TO MITOCHONDRIAL DNA DEPLETION: CLINICAL COURSE OF FOUR PATIENTS**

Danijela Petković Ramadža*, Tamara Žigman, Ruža Grizelj, Dorotea Ninković, Lana Omerza, Mina Natalija Anićić, Marijana Ćorić, Johannes A Mayr, René Feichtinger, Saskia Wirtmann, Holger Proksch, Ksenija Fumić, Jurka Vuković, Ivo Barić. University Hospital Centre Zagreb and

Mitochondrial DNA depletion syndromes (MDS) are a group of autosomal recessive disorders caused by disruption of mtDNA maintenance that results in reduced mtDNA content and disturbed energy production. MDS are genetically and phenotypically heterogeneous. One common phenotype is the hepatocerebral form that manifests in first months of life and causes early death due to liver failure. Liver transplantation (LTx) in not recommended in patients with neurological involvement. Our objective is to raise awareness about the clinical spectrum of early onset liver failure due to MDS.