Carnitine Uptake Defect (CUD)
(metabolic condition: fatty acid oxidation disorder)

Also known as:
• carnitine transport defect
• renal carnitine transport defect
• systemic carnitine deficiency

What are fatty acid oxidation disorders?
Fatty acid oxidation disorders (FAOD) are a group of inherited conditions in which fatty acids cannot be broken down to provide the body with energy. Fatty acids are a key source of energy during fasting or stress, such as during an illness. An inability to metabolize fatty acids leads to an accumulation of toxic metabolites in the body along with cellular energy deficiency which in combination can cause serious health problems.

What is CUD?
Carnitine is required to transport fatty acids into the mitochondria of the cell where breakdown occurs to produce energy. In CUD, carnitine that is present in the diet or synthesized by the liver cannot be transported into cells. This results in a deficiency of carnitine inside the cells leading to an inability to use fatty acids for energy production.

Maternal CUD is a condition that occurs when an infant is born to a woman with untreated CUD. In these situations, infants will have low carnitine at birth but are unaffected and do not require treatment.

What causes CUD?
CUD is caused by mutations in the carnitine transporter gene resulting in very low carnitine levels.

How common is CUD?
CUD is a rare condition with an estimated incidence of 1 in 50,000 infants born in the USA.

What are the clinical features of CUD?
Although infants with CUD may appear normal at birth, during a period of fasting (such as during a common illness), a child who was previously healthy in the first few months of life may present with lethargy, hypoketotic hypoglycemia and hypotonia. They may also present with cardiomyopathy, muscle weakness and an enlarged liver. They are at risk for encephalopathy, coma and sudden death.

What is the screening test for CUD?
A decreased concentration of carnitine is detected on the newborn blood spot screen. Newborn blood spot screening is not 100% perfect. Infants with clinical symptoms need timely assessment and diagnostic testing even if their screen result is normal.

How is the diagnosis confirmed?
The diagnosis of CUD is confirmed by the finding of very low total and free carnitine in a blood sample together with high levels of carnitine excreted in urine. Further testing may include enzyme analysis in a skin biopsy and/or mutation analysis of the carnitine transporter gene. Blood and urine carnitine levels in mothers of infants with low carnitine can distinguish between maternal CUD and CUD in the newborn. Specialists at the clinics listed below will arrange diagnostic testing.

How is CUD treated?
CUD is treated by supplementation with carnitine and avoidance of fasting. Parent education is an important component of treatment, including advising parents on actions to be taken when their child develops a minor illness. The treatment is lifelong.

What is the outcome of treatment for CUD?
Prevention of a metabolic crisis by carnitine supplementation and prevention of fasting is essential to ensure a good outcome for CUD. Prompt treatment of a metabolic crisis with intravenous fluids and glucose is necessary. Recurrent metabolic crises may result in neurological damage or death.

Is CUD inherited?
CUD is inherited as an autosomal recessive trait. Parents of a child with CUD are carriers of the condition and have a 1 in 4 chance of having another affected child in each subsequent pregnancy. CUD carriers are healthy. Genetic counselling and prenatal testing is available to all families with CUD.

For additional resources, please call:
Edmonton Medical Genetics Clinic
8-53 Medical Sciences Building
8440-112 St. N.W.
Edmonton, AB T6G 2H7
Phone: 780-407-7333
Fax: 780-407-6845
Emergency consultations:
Phone 780-407-8822 and ask for the genetic metabolic specialist on call.

Inherited Metabolic Disorders Clinic
Alberta Children’s Hospital
2888 Shaganappi Trail N.W.
Calgary, AB T3B 6A8
Phone: 403-955-7587
Fax: 403-955-3091
Emergency consultations:
Phone 403-955-7211 and ask for the specialist on call for metabolic diseases.

Early screening, detection and treatment – every infant, every time
For more information about the NMS Program, visit www.albertahealthservices.ca/newbornscreening.asp
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