DIAGNOSIS AND MANAGEMENT OF WILSON’S DISEASE IN CHILDREN: A TUNISIAN SINGLE-CENTER EXPERIENCE

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Introduction Wilson’s disease (WD) is an autosomal recessive genetic disease characterized by accumulation of copper in the body leading to severe hepatic and neurological damage. It is one of the few genetic diseases that can be treated effectively. The aim of the study was to describe Diagnostic features, therapeutic management and outcome of WD in Tunisian children.

Methods Retrospective cohort study of WD cases diagnosed at Tunisian tertiary referral hospital over a period of 10 years (2010 – 2019). Data collection was done from medical records.

Results We collected 20 cases of WD (15 boys and 5 girls). Median age at diagnosis was 8 years [3 - 12 years]. Consanguinity was found in 14 patients.

Three patients had a history of WD in the siblings. Clinical presentation was as following: four patients were diagnosed by family screening at a presymptomatic stage, 15 patients presented with hepatic symptoms (jaundice (n=4); hepatomegaly (n=5); clinical signs of portal hypertension (n=6)); and one patient presented with hematuria caused by kidney stones (hypercalciuria). Chronic hepatitis, acute hepatitis, liver cirrhosis and fulminant liver failure were observed respectively in two, four, six and four cases. Neurological and psychiatric involvement was noted at diagnosis in 7/20 patients. Kayser Fleischer’s ring was observed in three cases.

Proximal tubular involvement was reported in three cases, associated to glomerular involvement in one case. Hematologic disorders were observed in

11/20 cases: pancytopenia (n=2), thrombocytopenia (n=5), Coombs-negative hemolytic anemia (n=4). Thirteen patients had low levels of serum ceruloplasmin, with a median level of 0.12 g/L [0.04-0.18]. The 24-hours urinary copper test showed high level in 12 cases with a median level of 3.8 μmol/24h [1.5-6.72]. D-penicillamine sensitization test was performed in ten cases and was positive in eight cases. Liver biopsy was done in only two cases. Mutation analysis of the ATP7B gene was performed in four cases: no mutation was detected in one case and three patients had homozygous ATB7B mutation (H1069Q). D-penicillamine and pyridoxine were started in all cases.

One patient presented adverse reaction and received zinc acetate (Wilzin).

Most patients were stabilized or improved on chelation therapy, one patient deteriorated and one patient died within the follow up period.

Conclusion Cirrhosis at diagnosis increases the risk of death. Early diagnosis, at a precirrhotic stage was associated with a best prognosis.
prevalence of anemia was 90%. Only two patients received nutritional support.

**Conclusion** Malnutrition in the hospital is common in Tunisia but remains under diagnosed and insufficiently supported. This entity deserves further study to determine its impact on health expenditures and to improve its screening and management.

**HOW MANY CALORIES ARE ENOUGH?**
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Because of disease complexity, it is challenging to realistically estimate energy needs of cystic fibrosis (CF) patients. During regular dietician’s check-ups, we noticed that CF patients who gained weight or maintained BMI targets consume more calories than are estimated as enough, following the ECFS and ESPEN guidelines. Therefore, we question if 100% increase of recommended energy intake is enough for some patients.

To support this doubt we calculated patients’ energy needs based on Harris-Benedict formula and the level of physical activity, increased them following ESPEN guidelines and compared the recommended to actual intake and thriving.

Best example is a malnourished, picky-eater 12-year girl with poor appetite and impaired lung function (FEV1 29-43% p.v.). The girl finally agreed to PEG tube in order to increase feeding possibilities when the BMI was 13 kg/m2 (-3.03 SD) and FEV1 35%p.v. At that point, her estimated energy needs were 1900 kcal and according to ESPEN guidelines, the recommended intake was doubled. Because of poor food-intake (1000-1500 kcal), this was achieved mainly through enteral nutritional supplements either orally or via PEG. She still didn’t gain weight over a period of 9 months. During her last hospitalisation we reviewed the dietary approach. Her daily energy intake from food didn’t surpass 1300 kcal so we increased the enteral feeds to 5000 kcal daily thus mounting energy intake to = 6500 kcal/22.5 kg. After the lung function stabilized again (FEV1 37%p.v.), this high intake was necessary to retain weight gain. Following the same protocol at home, she gained 4 kg in a month (16.7% of weight at discharge). Her BMI is now 15.3 kg/m2 (-1.68 SD).

Our patient consumes 300-330% of energy needs of healthy peers. This example emphasizes the importance of the dietician in the CF team, an individual approach to each patient and ‘thinking out of the box’. Maybe it is time to revise the guidelines regarding energy requirements for CF patients.

**ACUTE ESOPHAGEAL NECROSIS IN A 15-YEAR OLD BOY – A CASE REPORT**
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Acute esophageal necrosis (black esophagus, Gurvits syndrome) is a rare clinical entity which leads to upper gastrointestinal bleeding. First description dates to 1990, with around 115 cases described in the literature. The condition has pathognomonic endoscopic appearance characterized by circumferential black mucosa in the distal esophagus, and discontinuing abruptly at the gastroesophageal junction. The pathogenesis is unclear, apparently multifactorial mucosal ischemia due to low flow vascular state or microvascular thrombosis is predisposing to topical damage by gastric content reflux. It’s commonly seen in elderly men, with risk factors like diabetes, malignancy, alcohol consumption, shock, major surgery.

Diagnosis is made endoscopically.

Management requires hemodynamic stabilization, acid suppressive medication with avoidance of nasogastric tube placement. The condition has very poor prognosis, with mortality rate up to 35%, and various complications including stenosis and perforation with mediastinitis and abscess formation.

Our patient, a 15 year old boy underwent surgery for scoliosis. During the immediate post surgical period he had hematemesis with consequent hemorrhagic shock. He was stabilized (IV fluids, packed red blood cells), nasogastric tube was inserted with evacuation of around 160 mL of blood and he was referred to our ICU. He required mechanical respiratory support and inotropic medications. Continuous parenteral PPI therapy was commenced.

Black, charcoal-like content was draining from the nasogastric tube, with further deterioration in hemoglobin levels.

Esophagogastroduodenoscopy showed black mucosa of lower esophagus, partly circumferential, partly linear, with cut-off at gastroesophageal junction.

There were no radiological signs of esophageal perforation, bilateral lung consolidates were surrounded by ground-glass interstitial changes.

Patient was kept NPO, on parenteral nutrition, with PPI and antibiotic treatment. He was weaned mechanical ventilation after three days, followed by brief stint of non-invasive respiratory support.

Unfortunately, significant stenosis with stricture formed in the area overlying initial necrosis. After several attempts of endoscopic balloon dilatation, refractory strictures reemerged. Surgical gastrostomy was performed to enable sufficient enteral caloric intake, and bring the patient to ideal physical condition for further treatment.

Planned colonic interposition surgery was not performed because of inadequate length of colon, hence thoracic surgeons performed retrosternal esophagogastropasty. Our patient had no further postoperative complications and was able to establish adequate oral feeding.

Acute esophageal necrosis should be considered as one of the causes of upper gastrointestinal bleeding, especially because its high mortality and complications rate requires immediate and aggressive early management.

**RISK FACTORS OF AUTOIMMUNE GASTRITIS IN CHILDREN WITH CELIAC DISEASE**

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The aim was to determine the risk factors of autoimmune gastritis (AG) in children with celiac disease (CD) Materials and