Abstracts

healthcare professionals should have greater awareness. This has been supported by the recent RCPCH position statement calling for practical sign-posting.7

G78 A SIMPLE SCREENING METHOD FOR THE MEASUREMENT OF LYSOosomal ACID LIPASE USING DRIED BLOOD SPOTS

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Background Deficiency of the enzyme lysosomal acid lipase (LAL) presents in infancy with hepatosplenomegaly, adrenal calcification and failure to thrive – Wolman Disease (WD). Cholesterol Ester Storage Disease (CESD) is the more common and milder form and presents later with hepatomegaly and hyperlipidaemia. Until recently there was no recognised method for the measurement of LAL in dried blood spots (DBS).

Aim To develop a simple method for the measurement of lysosomal acid lipase in DBS and validate the technique using samples from patients with CESD.

Methods LAL was extracted from DBS with water and incubated with β- pepstatin in acetate buffer and cardiolipin as an activator of LAL. Lalistat 2 is used as an inhibitor of LAL and activity is estimated by measuring total lipase activity and activity in the presence of Lalistat 2. LAL was determined by subtracting activity measured in the inhibited reaction from that in the uninhibited reaction.

The assay was performed using 96-well plates and read at 355nm/460nm.

Results LAL activity in normal controls was 0.37 – 2.30 nmol/punch/hr. Samples from CESD patients and WD have significantly reduced LAL activity: ≤ 0.04 nmol/punch/hr (n = 32). Activity in samples from obligate carriers is in the range 0.15 – 0.40 nmol/punch/hr (n = 15).

Conclusions Until now, methods for the measurement of LAL required the use of leucocytes and fibroblasts. These are difficult samples to prepare and methods for the measurement of LAL using these sample types are time consuming and expensive. Measurement of LAL in DBS is made difficult by the presence of other lipases in whole blood. Lalistat 2 is a specific inhibitor of LAL which allows the reliable determination of LAL in DBS. Results show the method differentiates clearly between normal controls, carriers and affected cases.

The DBS sample is simple to prepare and transport. By performing the assay using 96-well plates, samples can be processed efficiently in batches, previously not possible. Therefore it is now practical to screen for WD and CESD in patients with a lower index of suspicion of the disorder.

G79 NUTRITIONAL STATUS OF INFANTS WITH CONGENITAL HEART DISEASE

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Aims To assess the nutritional status of infants with congenital heart disease (CHD) who were managed jointly by the medical and dietetic teams and required either high calorie feeds or nasogastric feeding to optimise their growth.

Methods Infants with CHD requiring dietary input between 2003–2011 were reviewed. Patients were identified from paediatric cardiac and dietitians databases. A retrospective case note and electronic clinic letter review was performed.

Results Thirty nine infants were studied: 15(39%) with cyanotic CHD and 24(61%) acyanotic CHD. In 12 (29%) cases CHD was part of a syndrome. 25(64%) were treated surgically, 8(21%) required transcatheter procedures and 6(15%) were managed medically. Surgery was performed in 9 (27%) within 1-month of their birth and 24(73%) within the first-year.10 (67%) of the cyanotic infants had their corrective surgeries in the first 6 months. All 6 (15%) managed medically were among the acyanotic infants, 19 (76%) of the acyanotic infants were on diuretics and 10(40%) were operated in the later half of the year. There were 4 (10%) deaths in this cohort.

The birthweight was less than the 2nd centile in 12% of infants and overall 67% had a birthweight less than 50th centile. In the preoperative period, 14(41%) were nasogastric fed (NG). The majority of infants were on mixed feeds, 50% receiving some breast milk and 65% receiving Infatrini to maximise the weight gain. In the postoperative period, 10(30%) needed nasogastric feeds and 3(8%) with complex gut anomalies needed gastrostomy feeding tubes. The NG tubes were in place post-operatively for a median period of 2 months. At the end of 1 year, there was improvement in weight gain in 8(55%) of the cyanotic infants and 9(38%) of the acyanotic infants.

Conclusion Challenges persist in optimising the nutritional needs especially in children with CHD particularly those on prolonged diuretics. Collaboration with the dieticians is essential for optimising their growth prior to and after surgery.

G80 PRESENCE AND PROGRESSION OF IDIOPATHIC DILATED CARDIOMYOPATHY

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Aim Idiopathic Dilated cardiomyopathy (iDCM) is an uncommon presentation in children and young people but with significant and quite often poor prognosis. We aim to look at the presence and progression of iDCM.

Material and Methods 12 year retrospective analysis of data in a single centre; all patients with the term “DCM” were reviewed from the cardiac database (36 patient). A diagnosis was made on clinical and echocardiographic criteria. We excluded any patient where a cause was identified, analysed sequential echocardiograms and ECGs, and evaluated their progression and outcome.

Results There were 24 patients with iDCM; 15 boys and 9 girls. Presentation age range was Birth – 16 years years (mean: 5 years 8 months). Children/adolescents were followed up for an average of 80-months (range: 6 – 143 months). We measured LV dimensions and ventricular function and “z” scores calculated to track growth along with valvar lesions. 22 were on medication; b-blockers in 16, and 21 had an ACE inhibitor/ARB. 6 had additional medication in the form of Spironolactone and (12) diuretics. Anti-platelet (12)) use was also studied.

We had an adverse outcome (death/transplant) in 17%, death in 4 with 1 going on to have a transplant within this period. Of the survivors, we tracked their NYHA grade at the last review – (I, through IV). The fractional shortening was (7 – 32%) at presentation, (mean 18%), and 10 – 34% at discharge (mean 25%). Of those who died, the mean FS was 17%, and 28% in the survivors. LV diastolic dimension (LVDd, z scored) similarly was noted at presentation and at last review.

Conclusion We found that there was an adverse outcome in 17%. Actuarial survival was 100% at 1 year and 83% at 5 years with the best outcomes noted in younger age groups and in those where there was an improvement in systolic function.

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