# Training Syllabus CLINICAL SYLLABUS



## SYLLABUS FOR TRAINING IN CLINICAL PAEDIATRIC METABOLIC MEDICINE

Updated July 2006

This syllabus is intended as a guide. Whilst the training should be comprehensive, it is recognised that not all subjects can be covered in the same detail. Changes may also be necessary to meet local needs.

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Nutrition	Teaching	Academic/Research	
Genetics	Organisational	Managerial	
Development	Research		
Ethics			

## C.1 KNOWLEDGE

# C.1.1 Physiology and Biochemistry

The normal physiology and biochemistry, including changes during childhood, of:

- fluid and electrolyte balance.
- acid base regulation
- intermediary metabolism including blood glucose and metabolic response to fasting, lactate, ammonia, aminoacids, organic acids and fatty acids.
- oxidative phosphorylation and the respiratory chain
- lipids and lipoproteins
- cholesterol and other sterols
- lysosome and peroxisome metabolism
- purines and pyrimidines
- porphyrins
- calcium metabolism
- bilirubin
- trace metal metabolism
- relevant aspects of brain metabolism including neurotransmitters. The role of the blood brain barrier
- enzyme biochemistry and tissue expression

#### C.1.2 Paediatric Metabolic Disease

- The pathological and biochemical changes, clinical symptoms, investigations and management in metabolic disorders of these pathways and organelles, as listed in the appendix.
- Drug management. An understanding and experience of the drugs used for the treatment of metabolic disorders.
- Transplantation. The metabolic indications for and long term follow up of
  - 1. liver transplantation
  - 2. haematopoietic stem cell transplantation
  - 3. renal transplantation
- Principles of ex vivo and in vivo gene transfer and its relevance to IEM

#### C.1.3 Nutrition

- The normal nutritional requirements of protein, energy, vitamins and minerals for normal growth and development.
- The principles of dietary therapy including the consequences of changes in the intakes of nutrients. Consequences of under-nutrition and specific nutritional deficiencies.

#### C.1.4 Genetics

- Mechanisms of inheritance and an understanding of molecular genetics including mitochondrial DNA
- Understanding of principles of prenatal diagnosis

# C.1.5 Development

- Normal intellectual and psychological development
- Methods of assessment of DQ, IQ, behaviour and neuropsychological function

#### C.1.6 Research

• Principles of Good Clinical Practice (GCP) in Clinical Trials

# C.2. SKILLS

# C.2.1 Clinical skills

- 1. Assessment and investigation of metabolic causes of :
  - Acute encephalopathy
  - Neurological disease to include retardation, regression, fits, movements disorder, myopathy, etc
  - Liver disease including acute liver failure
  - Hepatosplenomegaly
  - Cardiomyopathy
  - Eye disease
  - Nutritional status and growth
  - Dysmorphic patients
  - Skeletal disorders
  - Skin disorders
  - Renal disorders including the Fanconi syndrome

- Muscle disease
- 2. Emergency management and indications for and the use of:
  - assisted ventilation
  - peritoneal dialysis
  - haemodialysis, haemofiltration and related techniques
  - intracranial pressure monitoring
- 3. Genetic counselling and counselling prior to prenatal diagnosis

## C.2.2 Technical Skills

- 1. Interpretation of the investigations including understanding analytical, physiological and nutritional factors that influence the results. The use of investigations for long term monitoring of metabolic disorders.
  - Aminoacids
  - Neurotransmitter and biogenic amine metabolites in CSF and other fluids
  - Organic acids (including interpretation of GC/MS data)
  - Ammonia
  - Intermediary metabolites: glucose, FFA, lactate, ketones
  - Carnitines
  - Enzymes studies: specific and non specific assays
  - Glycosaminoglycans
  - Lipids and lipoproteins
  - Peroxisomal enzymes and metabolites (VLCFA, pristanoate, phytanate, etc)
  - Purines and pyrimidines
  - Vitamin, mineral and other nutrient analyses including copper and caeruloplasmin
  - Cholesterol and other sterols
  - Porphyrins, bilirubin and related compounds
- 2. Indications and interpretation of:
  - fasting test for disorders of intermediary metabolism
  - allopurinol test
  - glucagon test
  - tests for co-factor responsive disorders (such as B12 test for MMA)
  - loading tests (protein, amino acid, MCT,LCT, glucose)
  - screening methods
- 3. Biopsies -indications for, planning and interpretation of:
  - skin biopsy (for fibroblast culture)
  - liver biopsy
  - muscle biopsy (needle and open)
  - and other biopsies
- 4. Experience of the use and interpretation of molecular genetic techniques
- 5. Practical aspects of dietary management:
  - evaluation of nutritional intake and requirements
  - low protein diets and those with controlled aminoacid intake (MSUD, PKU including the management in pregnancy)
  - low and very low fat diets
  - diets for peroxisomal disorders including adrenoleucodystrophy
  - galactose and fructose free diets
  - dietary management of glycogen storage disease
  - emergency /crisis regimens

- electrolyte replacement regimens (for treatment of Fanconi syndrome)
- 6. Interpretation of psychological and neuropsychometric testing

## C.2.3 Management skills

- Understanding of management skills required for development and use of resources in the metabolic unit, including budget control, contracting, strategic planning and writing a business plan.
- Experience of day to day running of a paediatric metabolic service including management of admission policies within the medical directorate system.
- Importance of and involvement in clinical audit
- structure and function of local, regional and national NHS organisation

## C.2.4 Academic skills

- Critical evaluation of clinical results from literature review and audit
- Manuscript preparation
- Oral presentation skills

# C.2.5 Teaching skills

- Formal and informal teaching skills at undergraduate and postgraduate level
- Development of core teaching material to enable participation in teaching programmes on a regular basis
- Organisation of a postgraduate teaching programme

## C.2.6 Organisational skills

 Routine departmental tasks including duty rotas, teaching programmes and postgraduate meetings

#### C.2.7 Research skills

- Design of clinical trials including medical statistics
- Data organisation and presentation
- Computer literacy

#### C.3. ATTITUDES

# C.3.1 Clinical

- Appreciate the patient and families understanding of metabolic disease
- Understanding of psychological stress of diagnosis of metabolic disorders
- Understanding of ethnic and cultural differences in attitudes to metabolic disorders.
- Understanding of the problems for the patient and family of progressive disorders and disorders in which the natural history is very difficult to predict such as Leigh's disease.
- Working in partnership with others who care for the children to include multidisciplinary teams, education authorities, etc.

## C.3.2 Communication

• Effective communication and appropriate approach to parents, children, colleagues GP and community health care staff, research and laboratory staff and managers

#### C.3.3 Education

• Commitment to continuing self-education and to teaching others

## C.3.4 Support and leadership

- Supportive and sympathetic approach to handling families which can be maintained under difficult circumstances
- Understanding and positive approach to the supervision of junior medical staff
- Recognition of and coping with stress in self and others
- Effective leadership of clinical teams

## C.3.5 Academic/Research

- Innovative attitude to clinical problems
- supportive to / active in research
- Collaborative attitude to local and national colleagues

#### C.3.6 Managerial

- Interest in the overall organisation of departmental activities
- Understanding approach to local management structure
- Constructive attitude to the process of decision making
- Acceptance of shared responsibility for the use of resources
- Ability to respond effectively to clinical complaints

## APPENDIX

#### INBORN ERRORS OF METABOLISM

- Disorders of aminoacid and peptide metabolism
- (Phenylketonuria including the management in pregnancy, homocystinuria, MSUD, etc)
- Disorders of organic acid metabolism
- (propionic acidaemia, methylmalonic acidaemia, etc)
- Hyperammonaemia and urea cycle disorders
- Disorders of carbohydrate metabolism
- (Glycogen storage disease, galactosaemia, fructosaemia, etc)
- Disorders of fatty acid oxidation (MCAD, LCHAD, etc)
- Disorders of ketone body metabolism
- Lysosomal storage disorders (MPS, etc)
- Disorders of lipoproteins and lipid metabolism
- Peroxisomal disorders (Adrenoleucodystrophy, Zellweger's syndrome, etc)
- Disorders of purine and pyrimidine metabolism
- Disorders of calcium metabolism
- Disorders of metal metabolism (Wilson's disease, Menkes syndrome, etc)
- Congenital lactic acidoses and metabolic myopathies
- Porphyrias
- Disorders of cholesterol, sterol and bile acid metabolism
- Disorders of vitamin metabolism (biotin, cobalamin, etc)
- Defects of membrane transport (lysinuric protein intolerance, etc)
- Defects of glycosylation
- Defects of connective tissue