

Second Congress and workshop on Inborn Error of Metabolic Program (Rare of the Rare)

27-28 Feb. and 1Mar , Tehran, Iran

Date:	Wed 27 Feb 2013
07:30-08:00	Registration
08:00-08:30	Opening
	Holy Quran
	Iranian National Anthem
08:30-09:00	Overview of workshop

Date:	Wed 27 Feb 2013
	Panel: Macromolecular Disease, Mitochondria, Peroxisom
09:00-09:30	All about the Mitochondrial Clinical (Dr. P. Karimzadeh)
09:30-10:00	All about the Mitochondrial Genetic (Dr. M. Houshmand)
10:00-10:30	Beta-Oxidation (Dr Mitra Nourbakhsh)
10:30-11:00	Break
11:00-11:30	Clinical approach to Fatty acid Oxidation defect (Dr. Rostampour)
11:30-12:00	CDG (Dr. P. Sarkhail)
12:00-12:30	Non Classical PKU (Dr. Setodeh)
12:30-13:30	Lunch & Pray
13:30-14:00	All about the Peroxisomal Clinical (Dr. Y. Shafeghati)
14:00-14:30	All about the Peroxisomal Genetic (Dr. K. Gaedi)
14:30-15:00	Case presentation (Dr. S. Akbaroghli)
15:00-15:30	Break
15:30-16:00	National Policy of IEM (Dr. Talebi)
16:00-16:30	Case Presentation (Dr. O. Aryani)
16:30-17:00	Case presentation

Date:	Thr 28 Feb 2013
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	Panel: Macromolecular Disease, Lysosom
09:00-10:00	lysosomal Storage Disease, Introduction Clinical and Biochemical (Dr. M. BaniKazemi)
10:00-10:30	Important Clinical point of lysosomal disorders (Dr. M.R. Ashrafi)
10:30-11:00	Break
11:00-11:30	Clinicogenetic approach of Mucopolysaccharidosis (Dr. A. Alaee)
11:30-12:00	Glycogen Storage Disease (Dr. Farshidi)
12:00-12:30	Sphingolipidosis (Dr. Sh. Salehpour)
12:30-13:30	Lunch & Pray
13:30-14:00	Orphan of the Orphan (Dr. O. Aryani)
14:00-14:30	Neurodegenerative with Brain Iron Accumulation (Dr. Shalbaf)
14:30-15:00	Nimann-Pick Disease A-B -C (Dr. Tonekaboni)
15:00-15:30	Break
15:30-16:00	Metabolic Cause of Epilepsy (Dr. Moravej)
16:00-16:30	Case Presentation (Dr. B. Bozorgmehr)
16:30-17:00	Case presentation (Dr. N. Almadani)

Date:	Fri 1 Mar 2013
	Panel: Micromolecular
09:00-09:30	Clinical and Biochemical analysis of PKU (Dr. Razaghi Azar)
09:30-10:00	Introduction to Aminoacidopathies (Dr. F. Rohani)
10:00-10:30	Genetic Diagnosis of Metabolic Disorders (Dr. Houshmand)
10:30-11:00	Break
11:00-11:30	Homocystinuria (Dr. Shakiba)
11:30-12:00	Introduction to the laboratory service on common metabolic disorders I (Dr. Mosalaei)
12:00-12:30	Overview of Organic Acidemia (Dr. Sohilipour)
12:30-13:30	Lunch & Pray
13:30-14:00	The role of Sonography in genetic disorder (Dr. Shahsavan)
14:00-14:30	Biochemistry or Genetic? Discussion (Dr. Houshmand, Dr. Mosalaei)
14:30-15:00	Case Presentation
15:00-15:30	Closing Ceremony