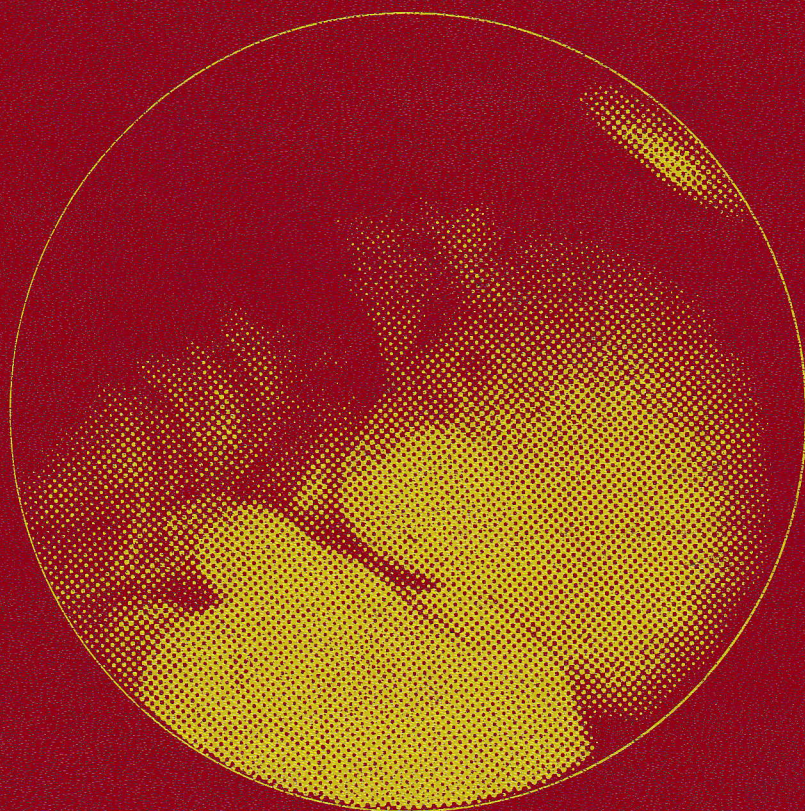


Hans Galjaard

GENETIC METABOLIC DISEASES

Early diagnosis
and prenatal analysis



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Contents

<i>Preface</i>	<i>ix</i>
<i>Acknowledgements</i>	<i>xi</i>
<i>Chapter I – Congenital Disease and (Infant) Mortality and Morbidity</i>	<i>1</i>
1. General introduction	3
2. Incidences and recurrence risks for various categories of congenital disorders	5
2.1. Introduction	5
2.2. Diseases due to single gene mutations	6
2.3. Chromosomal aberrations	9
2.4. Congenital malformations due to abnormal embryological development	12
3. Early diagnosis and prevention	15
3.1. Introduction	15
3.2. Analysis of metabolites	16
3.3. Demonstration of the responsible protein defect	18
3.4. Investigation of the gene mutation	21
3.5. Prenatal diagnosis and prevention	23
References	30
 <i>Chapter II – Diagnosis of Genetic Metabolic Diseases</i>	 <i>39</i>
1. General introduction to early diagnosis of genetic metabolic diseases	41
References	55
2. Inborn errors of carbohydrate metabolism	58
2.1. Introduction to differential diagnosis	58
2.2. Glycogenosis type II (Pompe's disease)	64
2.3. Glycogenosis type III (Cori's disease)	68
2.4. Glycogenosis type IV (amylopectinosis; Andersen's disease)	70

2.5. Transferase deficiency galactosemia	72
References	75
3. Mucopolysaccharidoses	82
3.1. Introduction to differential diagnosis	82
3.2. Mucopolysaccharidosis IH (Hurler's disease)	93
3.3. Mucopolysaccharidosis II (Hunter's disease)	101
3.4. Mucopolysaccharidosis III (Sanfilippo's syndrome)	105
3.5. Mucopolysaccharidosis IV (Morquio's syndrome)	109
3.6. Mucopolysaccharidosis IS (former type V of Scheie's syndrome)	112
3.7. Mucopolysaccharidosis VI (Maroteaux-Lamy syndrome)	115
3.8. Mucopolysaccharidosis VII	118
References	121
4. Diagnosis of mucolipidoses	132
4.1. Introduction to differential diagnosis	132
4.2. Mucolipidosis I (Sialidosis I)	135
4.3. Mucolipidosis II (I-cell disease)	137
4.4. Mucolipidosis III (pseudo-Hurler polydystrophy)	144
4.5. Mucolipidosis IV	146
4.6. α -Mannosidosis	147
4.7. Fucosidosis	151
References	158
5. Inborn errors in lipid metabolism	166
5.1. Introduction to differential diagnosis	166
References	174
5.2. Farber's lipogranulomatosis	178
References	181
5.3. Sphingomyelin lipidosis (Niemann-Pick disease)	183
References	191
5.4. Glucosylceramide lipidosis (Gaucher's disease)	195
References	202
5.5. Galactosylceramide lipidosis (globoid cell leukodystrophy or Krabbe's disease)	206
References	211
5.6. Sulfatide lipidosis or metachromatic leukodystrophy	215
5.7. Multiple sulfatase deficiency (mucosulfatidosis)	226
References	229
5.8. Ceramide trihexoside lipidosis (Anderson-Fabry disease)	237
References	246
5.9. GM ₁ -gangliosidosis	252
References	261
5.10. GM ₂ -gangliosidosis (Tay-Sachs disease and Sandhoff's disease)	266
References	281
5.11. Wolman's disease	290
References	294
5.12. Refsum's disease	297
References	300

6. Inborn errors of amino acid metabolism	303
6.1. Introduction to differential diagnosis	303
References	311
6.2. Inherited disorders in the metabolism of the urea cycle	315
6.3. Citrullinemia (argininosuccinate synthetase deficiency)	317
6.4. Argininosuccinic aciduria (argininosuccinate lyase deficiency)	320
6.5. Hyperargininemia (arginase deficiency)	323
References	325
6.6. Inherited disorders in the metabolism of branched-chain amino acids	330
6.7. Hypervalinemia and hyperleucine-isoleucinemia	333
6.8. Maple syrup urine disease	334
6.9. Isovaleric acidemia ("sweaty foot" syndrome)	340
6.10. β -Methylcrotonylglycinuria	343
6.11. α -Methylacetoacetyl-CoA β -ketothiolase (β -ketothiolase) deficiency	345
6.12. Propionic acidemia	347
6.13. Methylmalonic acidemia	351
References	355
6.14. Inherited disorders of sulfur amino acid metabolism	363
6.15. Homocystinuria	366
6.16. Cystathioninuria	379
6.17. Cystinosis	382
References	389
6.18. Defects in (hydroxy)lysine metabolism	401
6.19. Other inherited disorders of amino acid metabolism	407
References	411
7. Inborn errors in nucleic acid metabolism	415
7.1. Introduction to differential diagnosis	415
7.2. Lesch-Nyhan syndrome	422
7.3. Adenosine deaminase deficient combined immunodeficiency	436
7.4. Defective T-cell immunity (nucleoside phosphorylase deficiency)	444
7.5. Hereditary orotic aciduria	446
7.6. Xeroderma pigmentosum	448
7.7. Ataxia telangiectasia	457
References	459
8. Other genetic metabolic diseases	476
8.1. Introduction	476
8.2. Menkes' disease	477
8.3. Aspartylglucosaminuria	486
8.4. Familial hypercholesterolemia	491
References	497
<i>Chapter III – Methodology of Prenatal Analysis in Early Pregnancy</i>	505
1. General introduction	507
2. Amniocentesis in early pregnancy	512

3. Induction of second-trimester abortion	526
4. Composition and analysis of amniotic fluid supernatant	531
5. Alphafetoprotein assays and prenatal detection of open neural tube defects	535
6. Cultivation of amniotic fluid cells and fetal chromosome analysis	541
6.1. Introduction	541
6.2. Amniotic fluid cell cultivation	542
6.3. Different amniotic fluid cell types	546
6.4. Fetal chromosome preparation	548
6.5. Fetal sex determination	552
7. (Micro)biochemical analysis of cultured cells	554
7.1. General aspects	554
7.2. Preparation of the cell sample and incubation procedure	556
7.3. Microspectrophotometry and microfluorometry	568
7.4. Radiometric (micro)analysis	579
7.5. Electrophoresis, chromatography and other methods of prenatal analysis	586
7.6. Relation between cell cultivation conditions and biochemical properties	590
References	597
 <i>Chapter IV – Practical Experience with Prenatal Diagnosis of Genetic Metabolic Diseases</i>	 621
1. General introduction	623
2. Prenatal diagnosis of inborn errors in carbohydrate metabolism	630
3. Prenatal diagnosis of mucopolysaccharidoses and mucolipidoses	634
4. Prenatal diagnosis of lipidoses	641
4.1. Farber's disease	641
4.2. Niemann–Pick disease	642
4.3. Gaucher's disease	644
4.4. Krabbe's disease	645
4.5. Metachromatic leukodystrophy	646
4.6. Fabry's disease	649
4.7. GM1-gangliosidosis	650
4.8. GM2-gangliosidosis type I (Tay–Sachs disease)	652
4.9. Sandhoff's disease	655
4.10. Wolman's disease	655
5. Prenatal diagnosis of inherited aminoacidopathies	657
5.1. Citrullinemia	658
5.2. Argininosuccinic aciduria	658
5.3. Maple syrup urine disease	660
5.4. Propionic acidemia	661

	xix
5.5. Methylmalonic acidemia	662
5.6. Cystathioninuria	664
5.7. Homocystinuria	664
5.8. Cystinosis	665
6. Prenatal diagnosis of inherited disorders in nucleic acid metabolism and some other genetic defects	669
6.1. Introduction	669
6.2. Lesch-Nyhan syndrome	670
6.3. Xeroderma pigmentosum	673
6.4. Combined immune deficiency due to adenosine deaminase deficiency	676
6.5. Menkes' disease	678
6.6. Hypercholesterolemia	680
6.7. Lysosomal acid phosphatase deficiency	682
6.8. Hypophosphatasia	682
References	684
<i>Chapter V – Prevention of Sex-Linked Disease</i>	701
1. General aspects	703
2. X-linked metabolic disorders detectable by amniotic fluid cell analysis	710
2.1. Mucopolysaccharidosis II (Hunter's disease)	710
2.2. Ceramide trihexosidosis (Fabry's disease)	712
2.3. Lesch-Nyhan syndrome	713
2.4. Menkes' disease	715
3. Prevention by early diagnosis, genetic counseling and prenatal sex determination	717
3.1. Duchenne's progressive muscular dystrophy	719
3.2. Hemophilia	726
References	733
<i>Chapter VI – Some Aspects of Future Development</i>	743
1. Diagnosis and pathogenesis	745
2. Clinical treatment and basic research	751
3. Prevention and prenatal monitoring	759
References	778
<i>Appendices</i>	789
App. 1–10 – General methodology	791

App. 11–14 – Carbohydrate metabolism	809
App. 15–23 – Mucopolysaccharidoses	812
App. 24–40 – Mucolipidoses/lipidoses	817
App. 41–63 – Aminoacidopathies	830
App. 64–70 – Nucleic acid metabolism	847
App. 71–75 – Other genetic defects	853
<i>Subject index</i>	859