

3<sup>rd</sup> Edition

Johannes Zschocke  
Georg F. Hoffmann

# Vademecum Metabolicum

Diagnosis and Treatment  
of Inborn Errors of Metabolism

**milupa**  
METABOLICS

 Schattauer

## **Patterns of acute presentation**

<i>Main problem</i>	<i>See page</i>
The “metabolic emergency”	3
Early-onset seizures	14
Cardiomyopathy	17
Liver failure	19
Acute life-threatening episode, SIDS	24
Post-mortem investigations	25

## **Important laboratory findings**

<i>Main problem</i>	<i>See page</i>
Hypoglycaemia	5
Hyperammonaemia	7
Metabolic acidosis	10
Elevated lactate	12

Johannes Zschocke  
Georg F. Hoffmann

# **Vademecum Metabolicum**

---

3<sup>rd</sup> Edition

This page intentionally left blank

# Vademecum Metabolicum

**Diagnosis and Treatment  
of Inborn Errors of Metabolism**

---

3<sup>rd</sup>, revised Edition

**Johannes Zschocke, Innsbruck, Austria  
Georg F. Hoffmann, Heidelberg, Germany**

Foreword by  
William L. Nyhan, San Diego, USA

Contributors:  
Alberto B. Burlina, Padova, Italy  
Roberto Giugliani, Porto Alegre, Brazil  
Martin Lindner, Heidelberg, Germany  
Yoichi Matsubara, Sendai, Japan  
Ertan Mayatepek, Düsseldorf, Germany  
Shamima Rahman, London, UK  
Wolfgang Sperl, Salzburg, Austria  
Jerry Vockley, Pittsburgh, USA  
Kurt Widhalm, Vienna, Austria  
Ed Wraith, Manchester, UK

With 29 figures and 40 tables

**milupa**  
METABOLICS

 **Schattauer**

**Johannes Zschocke, Dr. med. habil., PhD**

Professor and Chair of Human Genetics  
Medical University Innsbruck  
Schöpfstraße 41, 6020 Innsbruck  
Austria

**Georg F. Hoffmann, Dr. med. habil.**

Professor and Chair of Paediatrics  
Ruprecht-Karls-University  
Im Neuenheimer Feld 430, 69120 Heidelberg  
Germany

This edition corresponds to the fourth German and Italian editions.

Bibliographic information published by the Deutsche Nationalbibliothek. The Deutsche Nationalbibliothek lists this publication in the Deutsche Nationalbibliografie; detailed bibliographic data is available in the Internet at <<http://dnb.d-nb.de>>.

**Important note:**

Medicine is an ever-changing science, so the contents of this publication, especially recommendations concerning diagnostic and therapeutic procedures, can only give an account of the knowledge at the time of publication. While utmost care has been taken to ensure that all specifications regarding drug selection and dosage and treatment options are accurate, readers are urged to review the product information sheet and any relevant material supplied by the manufacturer, and, in case of doubt, to consult a specialist. The publisher will appreciate – also in the public's interest – to be informed of possible inconsistencies. The ultimate responsibility for any diagnostic or therapeutic application lies with the reader.

No special reference is made to registered names, proprietary names, trade marks etc. in this publication. The appearance of a name without designation as proprietary does not imply that it is exempt from the relevant protective laws and regulations and therefore free for general use.

This publication is subject to copyright, all rights are reserved, whether the whole or part of the material is concerned. Any use of this publication outside the limits set by copyright legislation, without the prior written permission of the publisher, is liable to prosecution.

© 1999, 2004, 2011 by Milupa Metabolics GmbH & Co. KG, 61381 Friedrichsdorf, Germany  
[www.milupa-metabolics.info](http://www.milupa-metabolics.info)  
Printed in Germany

Composing: Bernd Burkart; [www.form-und-produktion.de](http://www.form-und-produktion.de)  
Printing and binding: CPI – Ebner&Spiegel, Ulm  
Printed on paper bleached without chlorine or acid.

ISBN 978-3-7945-2816-5

# Foreword

It is my pleasure to write the Foreword for the third edition of the *Vademecum Metabolicum*. This very useful book has now been translated into many languages including German, English, French, Italian, Hungarian, Portuguese and Japanese. It has continued to grow in scope, as has the field of inborn errors of metabolism. At the same time, it has remained true to its original objective of providing a systematic approach to the diagnosis of metabolic disease. The book is still small enough to fit in a pocket, and has become a favourite of physicians in training in Paediatrics and Genetics. Revisions have brought the book up to date, impressively so in the disorders of glycosylation, neurotransmission, and vitamin metabolism. Extensive tabular presentation leads the reader logically to the diagnostic possibilities. Optimal therapy, including dosages, makes for a well rounded approach to the various diagnosis and management of genetic diseases of metabolism.

University of California, San Diego, USA

**William L. Nyhan, MD, PhD**

# Preface

Inborn errors of metabolism, which cumulatively affect approximately one in every 500 newborns, represent a special challenge in general and paediatric practice. They frequently present with acute, life-threatening crises that require immediate specific intervention. The development and prognosis of the affected child may depend on rapid and effective treatment, but the large number of genetic defects in various biochemical pathways makes it difficult to be familiar with all diagnostic strategies and specific therapies. With this in mind, the *Vademecum Metabolicum* aims to provide practical guidance to the clinician.

This 3rd English edition has been completely revised and expanded. As in previous editions, the first section on the diagnosis and management of metabolic disorders includes clinical situations that may be caused by a metabolic disorder. Practical guidelines are discussed in detail and should reflect standard practice in many countries. The second section on individual metabolic pathways and their disorders has been completely revised and includes a considerable number of recently identified disorders. As in the previous editions, special emphasis has been placed on clinical features that are relevant to a whole group of diseases, useful diagnostic procedures (basic and special diagnostic tests) as well as details on emergency intervention and long-term treatment. The pathobiochemistry is described in more detail when it is relevant to the understanding of clinical symptoms and diagnostic tests. The sequence of the entries is either according to metabolic pathways or nomenclature.

The genetic basis of most disorders in the *Vademecum Metabolicum* has now been clarified, and the causative genes have been included when known. Throughout the text we have removed references to molecular studies as part of the diagnostic strategy since mutation analyses are now a standard option for confirmation of most metabolic disorders. Inheritance of the disorders is autosomal recessive unless specified otherwise.

We are grateful to Marinus Duran, Amsterdam, James V. Leonard, London, Verena Peters, Heidelberg, Jan A. M. Smeitink, Nijmegen, Udo Wendel, Düsseldorf, and Nicole Wolf, Amsterdam, who contributed to previous editions of this book. Again we are indebted to Dr. Beate Szczerbak, Milupa, Friedrichsdorf, for her unwavering, continuous support. The friendly and professional help of Claudia Ganter, Birgit Heyny and Klaus Jansch at Schattauer Publishing, Stuttgart, is gratefully acknowledged.

Innsbruck and Heidelberg, August 2011

**Johannes Zschocke**  
**Georg F. Hoffmann**



# Table of contents

<b>Diagnosis and management of metabolic disorders</b> . . . . .	1
<b>Essential basic laboratory tests</b> . . . . .	1
<b>General clinical situations</b> . . . . .	3
The metabolic emergency . . . . .	3
Hypoglycaemia . . . . .	5
Hyperammonaemia . . . . .	7
Metabolic acidosis and ketosis . . . . .	10
Elevated lactate . . . . .	12
Intellectual disability . . . . .	13
Metabolic (epileptic) encephalopathy . . . . .	14
The floppy infant . . . . .	16
Exercise intolerance . . . . .	16
Cardiomyopathy . . . . .	17
Dysmorphic features . . . . .	18
Liver disease . . . . .	19
Reye-like syndrome . . . . .	24
Sudden unexpected death (in infancy) . . . . .	24
Post-mortem investigations . . . . .	25
Fetal hydrops . . . . .	26
Unusual clinical observations . . . . .	27
Unusual laboratory findings . . . . .	28
Special metabolic investigations are not required in ... . . . .	29
<b>Special metabolic investigations</b> . . . . .	30
Simple metabolic urine tests . . . . .	30
Amino acids (AA) . . . . .	32
Organic acids (OA) . . . . .	33
Carnitine analyses . . . . .	34
Other special metabolic investigations . . . . .	35
Biopsies and enzyme studies . . . . .	38
Molecular genetic investigations . . . . .	39
<b>Function tests</b> . . . . .	41
Metabolic profiling . . . . .	41
Protein challenge . . . . .	42
Glucose challenge . . . . .	42
Prolonged fasting test . . . . .	43
Glucagon test . . . . .	45
Tetrahydrobiopterin (BH <sub>4</sub> ) test . . . . .	46
Phenylalanine loading test . . . . .	47
Allopurinol test . . . . .	48

<b>Newborn screening</b> . . . . .	49
Newborn screening for inborn errors of metabolism . . . . .	49
Newborn screening for non-metabolic disorders . . . . .	53
<b>Metabolic pathways and their disorders</b> . . . . .	54
<b>Amino acid and peptide metabolism</b> . . . . .	54
Principles of treatment . . . . .	55
Urea cycle disorders and inherited hyperammonaemias. . . . .	57
Organic acidurias . . . . .	61
Disorders of the metabolism of branched-chain amino acids not classified as organic aciduria . . . . .	66
Disorders of phenylalanine and tyrosine metabolism . . . . .	67
Disorders of the metabolism of sulphur amino acids . . . . .	69
Disorders of histidine, tryptophan and lysine metabolism. . . . .	72
Disorders of serine, glycine and glycerate metabolism . . . . .	73
Disorders of ornithine, proline and hydroxyproline metabolism . . . . .	75
Disorders of amino acid transport . . . . .	76
Other disorders of amino acid metabolism . . . . .	77
Disorders of the gamma-glutamyl cycle. . . . .	77
Disorders of peptide metabolism . . . . .	79
<b>Carbohydrate metabolism</b> . . . . .	80
Disorders of galactose and fructose metabolism . . . . .	81
Disorders of gluconeogenesis . . . . .	82
Glycogen storage diseases (GSD, glycogenoses) . . . . .	83
Disorders of glycerol metabolism . . . . .	86
Disorders of pentose/polyol metabolism . . . . .	86
Disorders of glucose transport . . . . .	87
Congenital hyperinsulinism (CHI). . . . .	88
<b>Fatty acid and ketone body metabolism</b> . . . . .	90
Disorders of fatty acid oxidation and ketogenesis. . . . .	91
Disorders of ketolysis . . . . .	95
<b>Energy metabolism.</b> . . . .	96
Disorders of pyruvate metabolism and the Krebs cycle . . . . .	96
Mitochondrial respiratory chain disorders. . . . .	99
Disorders of creatine biosynthesis. . . . .	111
<b>Purine and pyrimidine metabolism</b> . . . . .	112
Disorders of purine metabolism . . . . .	113
Disorders of pyrimidine metabolism . . . . .	115
Other disorders of nucleotide metabolism. . . . .	116
<b>Sterol metabolism</b> . . . . .	117
Disorders of sterol biosynthesis . . . . .	117
Disorders of bile acid synthesis . . . . .	120

<b>Porphyrin and haem metabolism</b> . . . . .	121
<b>Lipoprotein metabolism</b> . . . . .	124
Hypercholesterolaemias . . . . .	125
Hypertriglyceridaemias . . . . .	126
Mixed hyperlipidaemias . . . . .	126
Disorders of high density lipoprotein (HDL) metabolism . . . . .	127
Disorders with decreased LDL cholesterol and triglycerides . . . . .	128
<b>Lysosomal metabolism</b> . . . . .	129
Mucopolysaccharidoses (MPS) . . . . .	133
Oligosaccharidoses . . . . .	135
Sphingolipidoses . . . . .	136
Neuronal ceroid lipofuscinoses (NCL, CLN) . . . . .	140
Lysosomal export defects . . . . .	141
Other lysosomal disorders . . . . .	142
<b>Peroxisomal metabolism</b> . . . . .	143
<b>Protein glycosylation</b> . . . . .	146
Congenital disorders of glycosylation (CDG) . . . . .	146
<b>Neurotransmission</b> . . . . .	151
Disorders of biogenic amine metabolism . . . . .	151
Disorders of GABA metabolism . . . . .	154
Other neurometabolic disorders . . . . .	155
<b>Metabolism of vitamins and (non-protein) cofactors</b> . . . . .	156
Disorders of cobalamin absorption, transport and metabolism . . . . .	156
Disorders of folate metabolism and transport . . . . .	157
Disorders of biotin metabolism . . . . .	158
Disorders of pyridoxine metabolism . . . . .	159
Other disorders of vitamin metabolism . . . . .	161
<b>Metabolism of trace elements and metals</b> . . . . .	162
Disorders of copper metabolism . . . . .	162
Disorders of iron metabolism . . . . .	163
Disorders in the metabolism of other trace elements and metals . . . . .	164
<b>Other metabolic pathways</b> . . . . .	165
<b>Appendix</b> . . . . .	166
Helpful internet resources . . . . .	166
Free fatty acids and 3-hydroxybutyrate during fasting . . . . .	167
<b>Index</b> . . . . .	168



---

# Diagnosis and management of metabolic disorders

## Essential basic laboratory tests

The following basic laboratory tests should be performed in every child with an acute illness in whom a metabolic disorder is a possibility:

### *Blood glucose*

Hypoglycaemia is a presenting feature of several disorders particularly of carbohydrate and energy metabolism. Appropriate blood and urine samples should be obtained in the acute phase to make the correct diagnosis. *For details see page 5.*

### *Ammonia*

Ammonia is highly neurotoxic and hyperammonaemia carries a high but in principle avoidable mortality and morbidity. Urgent analysis of the plasma ammonia concentration is mandatory in all acutely ill neonates and all patients with undiagnosed encephalopathy. Facilities to determine ammonia at any time of the day should be available in all hospitals. Hyperammonaemia due to a primary urea cycle disorder is among the most urgent emergencies in metabolic paediatrics and will be missed if ammonia is not measured. *For details see page 7.*

### *Acid-base status*

Many metabolic disorders cause alterations in the acid-base status, both acidosis and alkalosis. Blood gas measurements need to be available at any time in every hospital. *For details see page 10.*

### *Lactate*

Elevated lactate is an important sequel of hypoxia and compromised energy metabolism and may be the cause of metabolic acidosis. A primary metabolic disorder should be considered if there is no convincing secondary cause such as shock, asphyxia or cardiac disease or in particular a difficult venepuncture. *For details see page 11.*

### *Urinary ketones (test strip)*

Ketonuria due to the ketone bodies 3-hydroxybutyrate and acetoacetate is normal during fasting but is pathological in the fed state and in the neonate where it may indicate a disorder of intermediary metabolism. Absence of ketones during fasting is suggestive of a fatty acid oxidation disorder. Ketone levels measured by non-specific tests (e.g. test strips) may be high due to the presence of interfering compounds. *See also page 11.*

### *Other laboratory tests*

Organ dysfunction caused by metabolic disorders may be recognised in routine investigations such as blood counts, liver function tests, coagulation studies or creatine kinase levels. Uric acid is elevated in several disorders with increased cellular turnover or decreased urinary clearance. *See also page 28.*

## Specific triggers of metabolic decompensation

<i>Triggers</i>	<i>Groups of disorders</i>
Vomiting, fasting, infection, fever, vaccination, surgery, accident/injury	Disorders of protein, energy or carbohydrate metabolism or hormone homeostasis
High protein intake and/or protein catabolism	Disorders of protein metabolism: aminoacidopathies, organic acidurias, urea cycle defects, hyperinsulinism-hyperammonaemia syndrome
Fruit, table sugar (sucrose), liquid medicines	Fructose intolerance
Lactose, milk products	Galactosaemia
High fat intake	Lipoprotein lipase deficiency, glycerol intolerance, fatty acid oxidation disorders
Drugs	Porphyrias, Glc-6-P-dehydrogenase deficiency
Extensive exercise	Disorders of fatty acid oxidation, glycolysis, muscle glycogenolysis, purine and pyrimidine metabolism, respiratory chain

## General clinical situations

### The metabolic emergency

**In the neonate**, the early clinical features of acute metabolic decompensation are almost always non-specific; they include “unwell”, lethargy, feeding problems, vomiting, abnormal breathing, hypotonia and seizures. Disorders of glucose, protein and fat breakdown (intermediary metabolism) in the neonatal period typically have an *asymptomatic interval*, with clinical manifestations from the second day of life onwards (“intoxication type”), although hyperammonaemia in particular may present as early as day 1. The baby’s general condition will usually deteriorate rapidly despite normal or non-specific findings in routine investigations (laboratory signs of infection, lumbar puncture, chest X-ray, cranial ultrasound) and antibiotic therapy. The *family history* may reveal siblings who died with similar clinical manifestations (“sepsis”, “SIDS”) or unexplained disorders in other family members (progressive neurological disease, maternal PKU, multiple miscarriages, HELLP syndrome, etc.). Consanguinity increases the risk of a recessive disorder. Metabolic disorders **after the neonatal period** may present with recurrent vomiting and lethargy progressing to coma without focal neurological signs or typical patterns of organ dysfunction. Initial management may follow similar principles as in neonates. Care must be taken to identify the conditions that triggered metabolic decompensation such as vomiting and fever or changes in the diet.

**A metabolic disorder should be considered**, along with other diagnoses (e.g. infection, CNS pathology) ...

- ... in all neonates with unexplained, overwhelming or progressive disease particularly after normal pregnancy and birth;
- ... in all children with acute deterioration of the general condition and/or reduced consciousness, particularly when preceded by vomiting, fever or fasting;
- ... in all children with symptoms and signs of acidosis or hypoglycaemia.

Appropriate diagnostic and therapeutic measures must be initiated as soon as possible to avoid long-term damage.

*Post-mortem investigations: see page 25*

### Phase 1: Basic metabolic emergency investigations and first line management

Stop intake of potentially toxic compounds (protein, fat, galactose, fructose)

*Insert i.v. line and take blood samples for urgent analysis of:*

- Electrolytes, *glucose*, CRP, CK, ALT, AST, creatinine, urea, uric acid, *acid-base status*, coagulation studies
- Ammonia, lactate
- Store plasma sample for amino acids, acylcarnitines, etc.
- Store filter paper card (“Guthrie” card for newborn screening) with dried blood spots for acylcarnitines (amino acids, possibly DNA studies)
- Store the rest of the other samples for possible additional tests (inform laboratory)

Download and install eVM - Vademecum Metabolicum 1.0.4 on Windows PC. The eVM is the electronic version of the Vademecum Metabolicum a highly successful...  
eVM - Vademecum Metabolicum on Windows Pc. Developed By: Johannes Zschocke. License: Free.