CORRECTION

Open Access

Correction to: Consensus guideline for the diagnosis and treatment of tetrahydrobiopterin (BH4) deficiencies



Thomas Opladen^{1*†}, Eduardo López-Laso^{2†}, Elisenda Cortès-Saladelafont^{3,4†}, Toni S. Pearson⁵, H. Serap Sivri⁶, Yilmaz Yildiz⁶, Birgit Assmann¹, Manju A. Kurian^{7,8}, Vincenzo Leuzzi⁹, Simon Heales¹⁰, Simon Pope¹⁰, Francesco Porta¹¹, Angeles García-Cazorla³, Tomáš Honzík¹², Roser Pons¹³, Luc Regal¹⁴, Helly Goez¹⁵, Rafael Artuch¹⁶, Georg F. Hoffmann¹, Gabriella Horvath¹⁷, Beat Thöny¹⁸, Sabine Scholl-Bürgi¹⁹, Alberto Burlina²⁰, Marcel M. Verbeek²¹, Mario Mastrangelo⁹, Jennifer Friedman²², Tessa Wassenberg¹⁴, Kathrin Jeltsch^{1†}, Jan Kulhánek^{12*†}, Oya Kuseyri Hübschmann^{1†} and on behalf of the International Working Group on Neurotransmitter related Disorders (iNTD)

Correction to: Orphanet Journal of Rare Diseases 15, 126 (2020)

https://doi.org/10.1186/s13023-020-01379-8

Following the original article's publication [1] the authors asked for the correction of Fig. 2, since the names of the disease genes [*GCH1* and *PCBD1*] in the figure published did not match the listed diseases [AR-GTPCHD and PCDD]. The correct Fig. 2 is shown below:

In the context of the manuscript correction and in order to match the text content, the words "apart from DHPRD" should be removed from the second row and second column of Table 4, as shown below:

The original article can be found online at https://doi.org/10.1186/s13023-020-01379-8.

* Correspondence: Thomas.Opladen@med.uni-heidelberg.de; Jan.Kulhanek@vfn.cz

[†]Thomas Opladen, Eduardo López-Laso, Elisenda Cortès-Saladelafont, Kathrin Jeltsch, Jan Kulhánek and Oya Kuseyri Hübschmann contributed equally to this work.

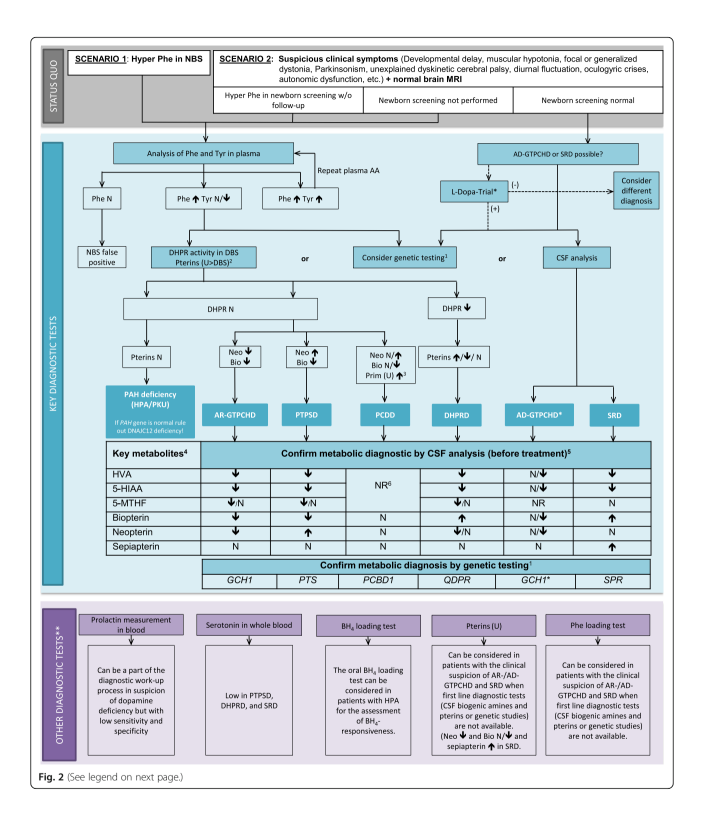
¹²Department of Paediatrics and Adolescent Medicine, First Faculty of Medicine, Charles University and General University Hospital in Prague, Prague, Czech Republic

Full list of author information is available at the end of the article



© The Author(s). 2020 **Open Access** This article is licensed under a Creative Commons Attribution 4.0 International License, which permits use, sharing, adaptation, distribution and reproduction in any medium or format, as long as you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons licence, and indicate if changes were made. The images or other third party material in this article are included in the article's Creative Commons licence, unless indicated otherwise in a credit line to the material. If material is not included in the article's Creative Commons licence and your intended use is not permitted by statutory regulation or exceeds the permitted use, you will need to obtain permission directly from the copyright holder. To view a copy of this licence, visit http://creativecommons.org/licenses/by/4.0/. The Creative Commons Public Domain Dedication waiver (http://creativecommons.org/publicdomain/zero/1.0/) applies to the data made available in this article, unless otherwise stated in a credit line to the data.

¹Division of Child Neurology and Metabolic Disorders, University Children's Hospital, Heidelberg, Germany



(See figure on previous page.)

Fig. 2 Diagnostic flowchart for differential diagnosis of BH₄Ds with and without HPA. ¹Consider genetic HPA workup depending on availability and financial resources. The gene panel should include the *QDPR*, *GCH1*, *PTS PCBD1*, *SPR* genes as well as *DNAJC12*. For *GCH1*, consider MLPA if Sanger sequencing is negative. ²The analysis in urine is more sensitive than in DBS and pathological patterns suggestive for PCDD and SRD can only be detected in urine but not in DBS. ³Primapterin measurement in urine is only elevated in PCDD. ⁴Aminoacids in CSF are not required for diagnosis of BH₄Ds. ⁵CSF analysis should always include standard measurements (cell count, proteins, glucose and lactate). ⁶Recommendation against measurements of HVA, 5-HIAA, 5-MTHF, and pterins in CSF in the case of PCDD. (*) A diagnostic L-Dopa trial should be limited to children with symptoms suggestive of dopa-responsive dystonia or to situations where biochemical and genetic diagnostic tools are not available. If the diagnostic L-Dopa trial is positive but the results of CSF biochemical and/or molecular genetic testing are not compatible with AD-GTPCHD or SRD, further aetiologies for dopa responsive dystonia should be considered (e.g. juvenile parkinsonism (PARK2gene)). (**) Can be considered if available. See text for more detailed information. Abbreviations: 5-HIAA, 5-hydroxyindoleacetic acid; 5-MTHF, 5-methyltetrahydrofolate; AA: amino acids; AD–/AR- GTPCHD: guanosine triphosphate cyclohydrolase I deficiency; BH₄, tetrahydrobiopterin; Bio: biopterin; CSF: cerebrospinal fluid; DBS: dry blood spot; DHPR: q-dihydropteridine reductase; DHPRD, dihydropteridine reductase deficiency; SRD: sepiapterin reductase deficiency; Tyr: tyrosine; u: urine; (+) = positive effect; (-) = no or no clear effect

Table 4 Recommended drugs and doses for BH_4 disorders

	Disorder	Starting dose	Doses	Target dose	Maximum dose	Management suggestion	Comment
irst line treatments							
Phe-reduced diet	All BH₄D with HPA					Titrate Phe restriction according to Phe levels in DBS or plasma	Follow PKU national treatment recommendations Use either Phe reduced diet or Sapropterin dihydrochloride to control Phe levels
Sapropterin dihydrochloride	All BH ₄ D with HPA	2-5 mg/kg BW/ day	Divided in 1–3 doses/ day	5–10 mg/kg BW/ day	20 mg/kg BW/day	Titrate dose according to Phe levels in DBS or plasma	Follow PKU national treatment recommendations Use either Phe reduced diet or Sapropterin dihydrochloride to control Phe levels
L-Dopa/DC inhibitor (carbidopa/ benserazide) 4:1	All BH₄D apart from PCDD	0.5 mg-1 mg/kg BW/day Dose recommendation relates to L-Dopa component!	Divided in 2–6 doses/ day	AD-GTPCHD: 3–7 mg/kg BW/ day All other BH ₄ D: 10 mg/kg BW/day or maximally tolerated dosage Dose recommendation relates to L-Dopa component!	Depending on clinical symptoms. Some patients need more than 10 mg/kg BW/day for resolving clinical symptoms	Increase 0.5–1 mg/kg BW/day per week Follow BW adaption until the BW of 40 kg. After 40 kg adjust depending on clinical symptoms Consider analysis of CSF HVA for dose adjustment	In young infants at least as many dosages as meals would be ideal (usually 5–6 /day)
5- Hydroxytryptophan (5-HTP)	All BH₄D apart from AD-GTPCHD and PCDD	1–2 mg/kg BW/ day	Divided in 3–6 doses/ day	Published target dose recommendations are highly variable 5-HTP doses are usually lower than L-Dopa doses		Titrate slowly (1–2 mg/kg BW/ day per week) depending on clinical picture and side effects Consider analysis of CSF 5HIAA for dose finding	5-HTP should follow L-Dopa/DCI treatment initiation Always in combination with a peripheral decarboxylase inhibitor (for example by simultaneous application with L-Dopa/DC inhibitor)
Folinic acid	In DHPRD and all BH₄D with Iow 5-MTHF in CSF		Divided in 1–2 doses/ day	10–20 mg/day		No titration needed Consider analysis of CSF 5MTHF for dose finding	
econd line treatments							
Pramipexole^a Dopamine agonist)	All BH₄D apart from PCDD	3.5-7 μg/kg/BW/ day (base) 5-10 μg/kgBW/ day (salt) Note: Distinction in salt and base content! (see product insert)	Divided in 3 equal doses/ day	Titrate to clinical Symptoms	75 μg/kg BW/day (3.3 mg/d base / 4 mg/d salt)	Increase every 7 days by 5 μg/kg BW/d	
Bromocriptine^a Dopamine agonist)	All BH₄D apart from PCDD	0.1 mg/kg BW/ day	Divided in 2–3 doses/ day	Titrate to clinical Symptoms	0.5 mg/kg/d (or 30 mg/d)	Increase every 7 days by 0.1 mg/kg BW/d	
Rotigotine ^a transdermal dopamine gonist)	All BH₄D apart from PCDD	2 mg/day		Titrate to clinical Symptoms	8 mg/day	Increase weekly by 1 mg	Children > 12 years Exchange patch every 24 h
Selegiline^a MAO B inhibitor)	All BH₄D apart from PCDD	0.1 mg/kg BW/ day	Divided in 2 (–3) doses/ day	Titrate to clinical Symptoms	0.3 mg/kg/d (or 10 mg/d)	Increase every 2 weeks by 0.1 mg/kg BW/d	Can cause sleep disturbances – morning and afternoon or lunchtime dosage is possible ATTENTION: orally disintegrating preparation needs much less dosage because the first-pass ef- fect of the liver is avoided

	Disorder	Starting dose	Doses	Target dose	Maximum dose	Management suggestion	Comment
Third line treatments							
Trihexyphenidyl ^a (Anticholinergic drugs)	All BH₄D apart from PCDD	< 15 kg: start 0.5– 1 mg/day > 15 kg: start 2 mg/day	< 15 kg: in 1 dose > 15 kg: in 2 doses	Effective dose highly variable (6–60 mg) Titrate to clinical Symptoms	Maximum dose: < 15 kg BW 30 mg/day > 15 kg BW 60 mg/d	Increase every 7 days by 1–2 mg/d in 2–4 doses/d	Consider side effects: like dry mouth, dry eyes, blurred vision (mydriasis), urine retention, constipation.
Entacapone ^a (COMT inhibitor)	All BH₄D apart from PCDD	200 mg (adult)			Up to 2.000 mg		In many countries licensed only for adults. Comedication with L-Dopa/DC inhibitor Consider reduction of concomitant L-Dopa supplementa- tion (10–30%)
Sertaline ^a (SSRI)	All BH₄D apart from PCDD	6–12 years: 25 mg/day in 1 dose > 12 years: 50 mg/day in 1 dose	6–12 years: in 1 dose > 12 years: in 1 dose	Children 50 mg/ day	50 mg/day < 12 years 200 mg/day > 12 years	6–12 years: increase after 7 days to 50 mg/ day in 1 dose > 12 years 50 mg/day in 1 dose	Don't stop treatment suddenly Note: Elevated risk of serotonin syndrome (SS) or malignant neuroleptic syndrome (MNS) when used with drugs impacting serotonergic pathway (e.g. 5-HTP, MAO inhibitors)
Melatonin ^a	All BH ₄ D apart from PCDD	0.01–0.03 mg/kg/ day			5–8 mg/day		Slow release preparation for sleep- maintenance insomnia available in some countries

Table 4 Recommended drugs and doses for BH₄ disorders (Continued)

Please note: The doses given are in a range typically used and have been published. In individual patients, some adjustment may be necessary depending on symptom response and side effects

^aThe evaluated literature did not provide BH₄D specific treatment dose recommendations for this drug. The listed doses, therefore, indicate treatment recommendations from Summary of Product Characteristics (SmPC) or neurotransmitter related publications (e.g. [119])

Abbreviations: 5-HIAA 5-hydroxyindoleacetic acid, 5-HTP 5-hydroxytryptophan, 5-MTHF 5- methyltetrahydrofolate, HVA Homovanillic acid, AD-GTPCHD Autosomaldominant guanosine triphosphate cyclohydrolase I deficiency, BH₂D Tetrahydrobiopterin deficiency, BW Body weight, COMT Catechol-O-methyl transferase, CSF Cerebrospinal fluid, DBS Dry blood spot, DC Decarboxylase, DCI Decarboxylase inhibitor, DHPRD Dihydropteridine reductase deficiency, L-Dopa L-3,4dihydroxyphenylalanine, MAO B Monoamine oxidase B, PCDD Pterin-4-alpha-carbinolamine dehydratase deficiency, Phe Phenylalanine, PKU Phenylketonuria, SSRI Selective serotonin reuptake inhibitor

Author details

¹Division of Child Neurology and Metabolic Disorders, University Children's Hospital, Heidelberg, Germany. ²Pediatric Neurology Unit, Department of Pediatrics, University Hospital Reina Sofía, IMIBIC and CIBERER, Córdoba, Spain. ³Inborn errors of metabolism Unit, Institut de Recerca Sant Joan de Déu and CIBERER-ISCIII, Barcelona, Spain. ⁴Unit of Pediatric Neurology and Metabolic Disorders, Department of Pediatrics, Hospital Germans Trias i Pujol, and Faculty of Medicine, Universitat Autònoma de Barcelona, Badalona, Spain. ⁵Department of Neurology, Washington University School of Medicine, St. Louis, USA. ⁶Department of Pediatrics, Section of Metabolism, Hacettepe University, Faculty of Medicine, 06100 Ankara, Turkey. ⁷Developmental Neurosciences, UCL Great Ormond Street-Institute of Child Health, London, UK. ⁸Department of Neurology, Great Ormond Street Hospital, London, UK. ⁹Unit of Child Neurology and Psychiatry, Department of Human Neuroscience, Sapienza University of Rome, Rome, Italy. ¹⁰Neurometabolic Unit, National Hospital, Queen Square, London, UK. ¹¹Department of Pediatrics, AOU Città della Salute e della Scienza, Torino, Italy. ¹²Department of Paediatrics and Adolescent Medicine, First Faculty of Medicine, Charles University and General University Hospital in Prague, Prague, Czech Republic. ¹³First Department of Pediatrics of the University of Athens, Aghia Sofia Hospital, Athens, Greece. ¹⁴Department of Pediatric, Pediatric Neurology and Metabolism Unit, UZ Brussel, Brussels, Belgium. ¹⁵Department of Pediatrics, University of Alberta Glenrose Rehabilitation Hospital, Edmonton, Canada. ¹⁶Clinical biochemistry department, Institut de Recerca Sant Joan de Déu,

CIBERER and MetabERN Hospital Sant Joan de Déu, Barcelona, Spain. ¹⁷Department of Pediatrics, Division of Biochemical Genetics, BC Children's Hospital, University of British Columbia, Vancouver, BC, Canada. ¹⁸Division of Metabolism, University Children's Hospital Zurich, Zürich, Switzerland. ¹⁹Clinic for Pediatrics I, Medical University of Innsbruck, Anichstr 35, Innsbruck, Austria. ²⁰U.O.C. Malattie Metaboliche Ereditarie, Dipartimento della Salute della Donna e del Bambino, Azienda Ospedaliera Universitaria di Padova -Campus Biomedico Pietro d'Abano, Padova, Italy. ²¹Departments of Neurology and Laboratory Medicine, Alzheimer Centre, Radboud University Medical Center, Donders Institute for Brain, Cognition and Behaviour, Nijmegen, The Netherlands. ²²UCSD Departments of Neuroscience and Pediatrics, Rady Children's Hospital Division of Neurology, Rady Children's Institute for Genomic Medicine, San Diego, USA.

Published online: 05 August 2020

Reference

 Opladen, et al. Consensus guideline for the diagnosis and treatment of tetrahydrobiopterin (BH4) deficiencies. Orphanet Journal of Rare Diseases. 2020;15:126. https://doi.org/10.1186/s13023-020-01379-8.