

Metagene: KNOWLEDGEBASE



Metagene

Knowledgebase for Diagnostic Support of Inborn Errors of Metabolism (IEM)

Metagene includes a diagnostic expert tool to identify possible diseases through laboratory values and symptoms.



Knowledgebase

Knowledgebase on the internet giving information about 1150 metabolic diseases and differential diagnoses.



DISEASE		SYMPTOM		LAB	
MIM	MIMGen	Disease	Synonym		
<input type="text"/>	<input type="text"/>	<input type="text"/>	<input type="text"/>		
202010	610613	11-BETA-HYDROXYLASE DEFICIENCY (CYP11B1)	ADRENAL HYPERPLASIA VI; STEROID 11 BETA-HYDROXYLASE DEFICIENCY		
264300	605573	17-BETA HYDROXYSTEROID DEHYDROGENASE III DEFICIENCY	PSEUDOHERMAPHRODITISM, MALE, WITH GYNECOMASTIA; HSD17B3		
616034	615787	2,4-DIENOYL-CoA REDUCTASE DEFICIENCY (DECRD)	2,4-DIENOYL-CoA REDUCTASE; DECRD; NADK2		
204750	614984	2-AMINOADIPIC ACIDURIA (AMOCAD, AMOXAD)	AMINOADIPIC ACIDURIA; AMOXAD		
600721	609186	2-HYDROXYGLUTARIC ACIDURIA (D) TYPE I	D-2-HYDROXYGLUTARIC ACIDURIA 1; D2HGA1		
613657	147650	2-HYDROXYGLUTARIC ACIDURIA (D) TYPE II	D-2-HYDROXYGLUTARIC ACIDURIA 2; D2HGA2		

In das Suchfeld DISEASE z.B. pku für Phenylketonuria eingeben

DISEASE		SYMPTOM	LAB
MIM	MIMGen	Disease	Synonym
<input type="text" value=""/>	<input type="text" value=""/>	<input type="text" value="pku"/>	<input type="text" value=""/>
261600	612349	PHENYLKETONURIA (PKU)	HYPERPHENYLALANINEMIA, MILD, INCLUDED; HPA, INCLUDED; PAH
261600		PHENYLKETONURIA, FETAL EFFECTS FROM MATERNAL (MPKU)	

PHENYLKETONURIA (PKU) für Anzeige der Details anklicken:

PHENYLKETONURIA (PKU)

Disease	PHENYLKETONURIA (PKU)
Synonym	HYPERPHENYLALANINEMIA, MILD, INCLUDED; HPA, INCLUDED; PAH
OMIM	261600 OMIM = Online Mendelian Inheritance of Men
Orphanet	79254
Protein (UniProt)	phenylalanine hydroxylase (PAH)
ExpASY	1.14.16.1
Gene locus	12q23.2 Detail information to gene locus by the National Center for Biotechnology Information NCBI: <ul style="list-style-type: none">• Phenylalanine hydroxylase
TreatableID	https://treatable-id.net
ICD	E70.0, E70.1

Laboratory findings:

1. **2-Hydroxyphenylacetic acid increased (urine)**
2. **3-Phenyllactic acid increased (urine)**
3. **Phenylacetic acid increased (urine)**
4. **Phenylalanine increased (plasma)**
5. **Phenylalanine/Tyrosine increased (plasma)**
6. **Phenylpyruvic acid increased (urine)**
7. **Phenylalanine increased (urine)**
8. 4-Hydroxyphenyllactic acid increased (urine)
9. 5-Hydroxyindolacetic acid (5-HIAA) normal/dec (cerebrospinal fluid)
10. Biopterin increased (urine)
11. D-Mannitol normal/inc (urine)
12. Ferric chloride reaction positive (urine)
13. Homovanillic acid (HVA) normal/dec (cerebrospinal fluid)
14. L-Tyrosine decreased (urine)
15. N-Acetylphenylalanine increased (urine)
16. Neopterin increased (urine)
17. Phenylalanine increased (cerebrospinal fluid)

Symptoms:

1. **intellectual disability/intellectual developmental disorder**
2. **musty body odor**
3. **blue eyes**
4. **fair hair**
5. **behavior, autism or autistic-like**
6. **decreased body height**
7. embryopathy
8. **hypertonia, spasticity**
9. hypopigmentation
10. **irritability**
11. **low birthweight (small for gestational age)**
12. **microcephaly (<2 SD for age)**
13. pigmentation, skin and sclera
14. seizures
15. **skin rash, eczematous or seborrheic**
16. **unusual odor / odour**
17. **vomiting**
18. Amino acids, plasma
19. behavior, hyperactive, restless
20. behavior, self-mutilating or destructive
21. childhood
22. EEG abnormalities [-]
23. epilepsy
24. infancy
25. leukodystrophy
26. mental retardation
27. MRI, brain, white matter abnormalities [-]
28. newborn, neonatal
29. Organic acids, urine
30. sleep disturbances
31. small for gestational age (SGA), intrauterine growth retardation (IUGR)

Literature:

1. Autor: Susan E Waisbren, (2025)
[Beyond neuropsychological tests: AI speech analysis in PKU](#)
2. Autor: François Feillet, (2025)
[Long-term safety of sapropterin in paediatric and adult individuals with phenylalanine hydroxylase deficiency: Final results of the Kuvan® Adult Maternal Paediatric European Registry multinational observational study](#)
3. Autor: Ahring K, (2024)
[Management of phenylketonuria in European PKU centres remains heterogeneous](#)
4. Autor: Feillet F, (2024)
[Efficacy and safety of sapropterin before and during pregnancy: Final analysis of the Kuvan® Adult Maternal Paediatric European Registry \(KAMPER\) maternal and Phenylketonuria Developmental Outcomes and Safety \(PKUDOS\) PKU-MOMs sub-registries](#)
5. Autor: Trefz F, (2024)
[Does hyperphenylalaninemia induce brain glucose hypometabolism? Cerebral spinal fluid findings in treated adult phenylketonuric patients](#)
6. Autor: Cary O Harding, (2024)
[Sepiapterin: a potential new therapy for phenylketonuria](#)
7. Autor: Wendy E Smith, (2024)
[Phenylalanine hydroxylase deficiency diagnosis and management: A 2023 evidence-based clinical guideline of the American College of Medical Genetics and Genomics \(ACMG\)](#)
8. Autor: WEBSITE MANAGER: Nenad Blau (2023)
[BioPKU](#)
9. Autor: Alghamdi MA, (2023)
[Classical phenylketonuria presenting as maternal PKU syndrome in the offspring of an intellectually normal woman](#)
10. Autor: Chen A, (2023)
[Clinical, genetic, and experimental research of hyperphenylalaninemia](#)
11. Autor: Cannet C, (2023)
[Phenylketonuria \(PKU\) Urinary Metabolomic Phenotype Is Defined by Genotype and Metabolite Imbalance: Results in 51 Early Treated Patients Using Ex Vivo 1H-NMR Analysis](#)
12. Autor: Muri R, (2022)
[Cortical thickness and its relationship to cognitive performance and metabolic control in adults with phenylketonuria](#)
13. Autor: Dobrowolski SF, (2021)
[Phenylalanine hydroxylase deficient phenylketonuria comparative metabolomics identifies energy pathway disruption and oxidative stress](#)
14. Autor: Dobrowolski SF, (2021)
[Mesenchymal stem cell energy deficit and oxidative stress contribute to osteopenia in the Pah enu2 classical PKU mouse](#)
15. Autor: Evers RAF, (2020)
[Tetrahydrobiopterin treatment in phenylketonuria: A repurposing approach](#)
16. Autor: Krämer J (2020)
[Case-control study about the acceptance of Pegvaliase in Phenylketonuria](#)

Metagene: DIAGNOSE TOOL



reset next

 possible diseases: 180

- ∨ CARDIOVASCULAR SYSTEM
 - └ GENERAL
- ∨ DIGESTIVE SYSTEM
 - └ GENERAL
- ∨ LABORATORY ABNORMALITIES
 - └ ROUTINE LAB, POCT

Select at least one organ system with abnormal clinical findings. Click next Q X

- ∨ CARDIOVASCULAR SYSTEM
 - CONGENITAL HEART DISEASE
 - GENERAL
- ∨ CHEST, BREAST, BACK, ABDOMEN
 - GENERAL
- ∨ DIGESTIVE SYSTEM
 - GENERAL
 - LIVER/GALLBLADDER, PANCREAS, SPLEEN
 - NUTRITION
- ∨ ENDOCRINE, NUTRITIONAL AND METABOLIC DISEASES
- ∨ LABORATORY ABNORMALITIES
 - ROUTINE LAB, POCT

Mindestens ein Organsystem oder Labormethode wählen, z.B. CARDIOVASCULAR SYSTEM für arrhythmia, DIGESTIVE SYSTEM für vomiting, ROUTINE LAB für hypoglycemia. Klick auf **“next”**

back next

Metagene
possible diseases: 9

cardiac arrhythmia, dysrhythmia

hypoglycemia

vomiting

Q X

hypoglycemia

hypoketotic hypoglycemia

Über das Suchfeld klinische Symptome oder Laborbefunde nacheinander suchen und auswählen. Klick auf **“next”**.

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ICD	Disease	MIM
E71.1	3-HYDROXY-3-METHYLGLUTARYL-CoA LYASE DEFICIENCY (HMGCLD)	246450
E71.1	BARTH SYNDROME (BTHS)	302060
E71.3	CARNITINE PALMITOYL TRANSFERASE DEFICIENCY (III), CPT2, MYOPATHIC, STRESS-INDUCED	255110
E71.3	CARNITINE-ACYLCARNITINE TRANSLOCASE DEFICIENCY (CACT)	212138
H49.8	KEARNS-SAYRE SYNDROME (KSS)	530000
E71.3	LONG-CHAIN-3-HYDROXYACYL-CoA DEHYDROGENASE DEFICIENCY (LCHAD)	609016
E71.3	MEDIUM CHAIN ACYL-CoA DEHYDROGENASE DEFICIENCY (MCAD)	201450
E71.1	PROPIONIC ACIDEMIA (PA, PCCA)	606054
E71.1	PROPIONIC ACIDEMIA (PA, PCCB)	606054

9 Krankheiten mit cardiac arrhythmia, vomiting und hypoglycemia werden gelistet.