

# Order form for molecular genetic testing

Zentrum Medizinische Genetik Würzburg, Biozentrum, Am Hubland, 97074 Würzburg



## Patient details (use label):

male  female

Given name \_\_\_\_\_

Name \_\_\_\_\_

Date of birth \_\_\_\_\_

## Cost coverage declaration:

- Invoice to patient (give details and address on reverse page)  
 Invoice to referring institution  
 Invoice by special agreement with Zentrum Medizinische Genetik Würzburg (to be obtained prior to testing)

## Relevant clinical patient and/or family data (please enclose reports, if available):

- Clinical diagnostics  
 Predictive / prenatal / carrier diagnostics  
 Known Index patient / known mutation:  yes  no

Gene: \_\_\_\_\_ Mutation: \_\_\_\_\_

## Test order (please tick boxes)

Please note the separate test orders for panel diagnostics on our homepage:  
<https://www.biozentrum.uni-wuerzburg.de/humangenetik/patientenversorgung/formulare/>

**MATERIAL: 5-10 ml EDTA- blood.** Please label tubes clearly, package in shatter-resistant packaging; ship at ambient temperature as soon as possible within one week

### Neuro-muscular disorders

- DMD/BMD (del/dup), (MLPA)  
 DMD/BMD (point mutations)  
 Emery-Dreifuss muscular dystrophy (aut. dom.)  
 Emery-Dreifuss muscular dystrophy (X-linked)  
 Facio-scapulo-humeral MD 1 (FSHD1, D4Z4)  
 Facio-scapulo-humeral MD 2 (FSHD2, SMCHD1)  
 Muscle hypertrophy (Myostatin)\*  
 Oculo-pharyngeal muscular dystrophy (OPMD)  
 Rigid Spine muscular dystrophy (SEPN1)  
 Spinal muscular atrophies (5q-associated)  
 Spino-bulbar muscular atrophy (Kennedy)

### Myotubular myopathies

- BIN1 (autosomal recessive)  
 DNM2 (autosomal dominant)  
 MTM1 (X-linked)

### Myofibrillar myopathies (MFM)

- BAG3  FLNC  
 CRYAB  MYOT  
 DES  TTN (Hot spots)\*  
 DNAJB6\*  LDB3 (ZASP)  
 FHL1

### Distal Myopathies

- DES  FLNC  
 DNAJB6\*  MYH7

### Myotonic dystrophies

- DM1 (Curschmann-Steinert)  
 DM2 (Proximal myotonic myopathy)

### Structural myopathies – Malignant Hyperth.

- Central core disease ( RYR1)  
 Malignant hyperthermia (RYR1/CACNA1S)  
 Multi minicore disease (SEPN1/RYR1)  
 Nemaline myopathy (ACTA1)

### Limb girdle muscular dystrophies

- Type EDMD (LMNA)  
 Type RMD (CAV3)  
 Type LGMD R1 (CAPN3)  
 Type LGMD R2 (DYSF)  
 Type LGMD R3 (SGCA)  
 Type LGMD R4 (SGCB)  
 Type LGMD R7 (TCAP)\*  
 Type LGMD R9 (FKRP)  
 Type LGMD R10 (TTN, Hot spots)  
 Type LGMD R12 (ANO5)

**If you want to order more myopathy genes tests, please use the order form for Myopathy Panel**

### Neuro-degenerative disorders

- Chorea Huntington (HTT)

### Coagulation disorders

- Hemophilia A (F8)  
 Hemophilia B (F9)  
 Hereditary angioedema type I & II (Serping1)

### Hereditary hearing loss

- CX26 (GJB2)\*,  CX30 (GJB6)\*  
 STRC\*

### Craniosynostoses

The order form for the Craniosynostoses can be found on our homepage

### FGFR3 associated skeletal dysplasias

- Achondroplasia (FGFR3)  
 Hypochondroplasia (FGFR3)  
 Thanatophoric Dysplasia (FGFR3)

### Limb malformations

- Split hand/foot\*  SHFM3\*,  SHFLD3\*  
 TAR-Syndrome (Del 1q21.1; RBM8A)\*

### Others

- Alkaptonuria (HGD)\*  
 CMT 1A/HNPP, (PMP22, MLPA)  
 Hypophosphatasia (ALPL)  
 Short stature (SHOX)  
 Lipodystrophy (LMNB2)\*  
 Marfan syndrome (FBN1)  
 Micro-deletion screening (MLPA)  
 Neurofibromatosis (NF1)  
 Polycystic kidney disease (PKD1, PKD2)  
 Pyruvate kinase deficiency (PKLR)

\* not accredited

Sample taken (date): \_\_\_\_\_ by: \_\_\_\_\_

Doctor's name (please print) \_\_\_\_\_

Doctor's signature \_\_\_\_\_

(Doctor's stamp)

According to the German gene testing act written patient's consent is required for every genetic test  
<https://www.biozentrum.uni-wuerzburg.de/humangenetik/patientenversorgung/forms-en/>