

# Metabolic Myopathies

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**METABOLIC MYOPATHIES** - decreased muscle energy supply due to biochemical abnormalities.

CARBOHYDRATES are essential for *anaerobic* energy needs (primarily – **cytoplasmic GLYCOGEN** → glycogenolysis → glycolysis).  
LIPIDS are essential for *aerobic* energy needs during sustained exercise (primarily - **serum LONG-CHAIN FATTY ACIDS** → β-oxidation in mitochondria).

1. **DYNAMIC (exercise-induced) myopathies** - symptoms (acute myalgias, stiffness → contractures, intermittent weakness → myoglobinuria\*) appear during / after exercise:
    - \* drink fluids after exercise!
    - A) **carbohydrate metabolism disorders** - **type V** (most common), **type VII-XI glycogenoses, Satoyoshi disease**; see p. 734-738 >>
      - *hemolytic anemia* accompanies only type VII (mild) and type IX (severe).
    - B) **lipid metabolism disorders** - **carnitine palmitoyl transferase deficiencies** see p. 750 >>
    - C) **purine metabolism disorders** - **myoadenylate deaminase deficiency** see below >>
    - D) **mitochondrial myopathies** - **succinate dehydrogenase deficiency**
  - *exercise intolerance* in childhood;  
*exertion-induced symptoms* (muscle pain, weakness, myoglobinuria) in 2-3<sup>rd</sup> decade.
    - **contractures** cause intense muscle pain, are electrically silent and not associated with ATP depletion.
    - exercise tolerance can be enhanced by slow induction phase (warm-up) or brief rest periods allowing for start of "*second-wind*" *phenomenon* (i.e. patient can continue exercise at previous level of activity after brief rest - switching to utilization of fatty acids).
  - *between attacks*, muscle strength, diagnostic test results are normal (may become abnormal with advancing age).
2. **STATIC (stable or slowly progressive) myopathies** - chronic fixed progressive weakness (simulates muscular dystrophy; no exercise intolerance, no myoglobinuria):
    - A) **carbohydrate metabolism disorders** - **type II-IV glycogenoses**. see p. 734-738 >>
    - B) **lipid metabolism disorders** - **carnitine deficiencies**: see p. 750 >>
      - 1) primary (muscle / systemic)
      - 2) secondary (β-oxidation defects, valproic acid)
    - C) **mitochondrial myopathies** (most)

N.B. type I and VI glycogenoses do not affect muscles!

## DIAGNOSIS

1. **Forearm (grip) exercise** - information about glycolytic (anaerobic) metabolism by evaluating *lactate* production in **ischemic exercise**:
  - rested, rested and fasting patient **repetitively squeezes handheld ergometer** while BP cuff is maintained above systolic pressure (induced ischemia prevents oxidative phosphorylation).
    - a) workload 4-7 kg-m at 60 Hz for 1 min (such duration does not induce ischemic pain).
    - b) sustain 1.5-second contractions separated by 0.5-second rest periods for 1 minute.
    - c) squeeze to 50% of maximum grip strength until exhaustion (usually ≈ 10 minutes).
  - **nonischemic workload** > 6-7 kg-m (well exceeds aerobic threshold) also produces comparable results and *avoids induced ischemia* (may cause severe muscle necrosis in glycolytic defects).
  - venous [lactate] and [ammonia] are determined from antecubital vein proximal to deep veins of forearm (e.g. median vein):
    - pre-exercise;
    - postexercise (1, 2, 4, 6, 10 minutes).
  - **normally**: [lactate] rises 3-5-fold within 1-2 minutes after exercise;  
[ammonia] rises 2-10-fold within 2-5 minutes after exercise.
    - glycogenesis* – [lactate] elevation does not occur (or is diminished); muscle develops painful contracture;
    - lipid metabolism disorders* – normal profile;
    - myoadenylate deaminase deficiency* – [ammonia] elevation does not occur;
    - mitochondrial disorders* – excessive [lactate] elevation;
    - poor effort* – neither [lactate] nor [ammonia] increase.
2. **Incremental bicycle ergometry** - information about aerobic metabolism.
3. **<sup>31</sup>P MR spectroscopy** - information about intracellular energy metabolites (i.e. ATP, inorganic phosphate, phosphocreatine).
4. **EMG**:
  - A) DYNAMIC myopathies:
    - *during episode* - electrical silence.
    - *after episodes* of severe myoglobinuria - myopathy and fibrillations.
    - *between episodes* – normal.
  - B) STATIC myopathies – myopathy, excessive irritability (incl. myotonic discharges, particularly in lumbosacral paraspinal muscles in Pompe disease).
5. **Muscle biopsy**:
  - 1) scattered **necrotic & regenerating fibers** (esp. after rhabdomyolysis episode).
  - 2) **specific findings** (e.g. vacuolar glycogen or lipid accumulations).
  - 3) specific **enzyme deficiency** (alternatively skin fibroblasts, intestinal mucosa, lymphocytes may be examined) – definitive diagnosis!
6. **Serum CK** moderately increased (very increased after attacks\* and usually normal between attacks of DYNAMIC myopathies).
 

\*together with myoglobinuria
7. **Genetic analysis for mutations**

## MYOADENYLATE DEAMINASE DEFICIENCY

**Myoadenylate deaminase (s. muscle AMP deaminase)** provides short-term ATP supply by catalyzing conversion of AMP → IMP through removal of ammonia. see p. 832 >>

- a) exertional myalgia ± myoglobinuria (**DYNAMIC myopathy**)
  - b) **asymptomatic** (myoadenylate deaminase gene 1p13-21 is mutated in ≈ 2% normal people).
- **forearm exercise test** - no increase in [ammonia].

BIBLIOGRAPHY for ch. "Metabolic Disorders" → follow this [LINK >>](#)

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**Viktor's Notes<sup>SM</sup> for the Neurosurgery Resident**  
Please visit website at [www.NeurosurgeryResident.net](http://www.NeurosurgeryResident.net)