Anesthesia recommendations for patients suffering from

Merosin-deficient congenital muscular dystrophy

**Disease name:** Merosin-deficient congenital muscular dystrophy

**ICD 10:** G71.0

**Synonyms:** Complete merosin deficient congenital muscular dystrophy, Congenital muscular dystrophy with primary laminin 2 (merosin) deficiency, Merosin negative congenital muscular dystrophy, LAMA-2 related muscular dystrophy (early and late onset LAMA-2 deficiency), Congenital muscular dystrophy, type 1A, MDCA1A

Congenital muscular dystrophies (CMDs) are rare, autosomal recessive genetic disorders that are clinically and genetically heterogeneous. All CMDs are characterized by progressive muscle weakness, delayed motor development, and dystrophic changes on muscle biopsy.

Merosin-deficient-congenital muscular dystrophy (MD-CMD) is characterized by severe progressive muscle weakness that results in contractures, scoliosis, and restrictive pulmonary disease. MD-CMD is the most common and severe form, representing 40% of all CMDs. It is caused by a mutation in the laminin α2 gene (LAMA2-on chromosome 6q22-23) resulting in absence of the laminin α2 chain (aka merosin) around muscle fibres. In addition to the clinical characteristics described above, patients with MD-CMD may present with elevated creatinine kinase (CK) diffuse white matter hyperintensities on brain MRI and seizures (in 30% of patients). Most of these patients have normal intelligence. Approximately one third of MD-CMD patients also have cardiac abnormalities including arrhythmias, and dilated cardiomyopathy. Anesthesia for these patients is required for both diagnostic and surgical procedures. Typical diagnostic procedures include MRI, endoscopies and muscle biopsies. Typical surgical procedures include contracture release, spine fusion, tracheostomy, and gastrostomy.

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**Medicine in progress**

- Perhaps new knowledge
- Every patient is unique
- Perhaps the diagnostic is wrong

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Find more information on the disease, its centres of reference and patient organisations on Orphanet: [www.orpha.net](http://www.orpha.net)
Disease summary

The main anesthetic concerns include:

1. Difficult airway – due to hypotonia, limited mouth opening from jaw contractures, maxillary discrepancy and macroglossia
2. Intra- and post-operative management of respiratory function, which is frequently compromised
3. Use of depolarizing relaxants – due to potential hyperkalemic cardiac arrest and rhabdomyolysis (controversy still exists around the use of halogenated agents; our preference is to avoid these agents)
4. Cardiovascular instability in patients with cardiac compromise
5. Hypoglycemia

Typical surgery

Surgical procedures:
Tracheostomy, gastrostomy tube placement, orthopedic procedures (i.e. posterior spinal fusion and tendon releases for treatment of muscle contractures), maxillofacial procedures and dental care

Diagnostic procedures:
Esophagoscopy and muscle biopsy.

Type of anaesthesia

There is no absolute contraindication for either general anesthesia or regional anesthesia.

The main anesthetic concern with general anesthesia is the use of depolarizing relaxants due to potential hyperkalemic cardiac arrest and rhabdomyolysis; controversy still exists around the use of halogenated agents. There are only three reports in the literature (six patients total) of anesthetic management in patients with MD-CMD. Every report used nontriggering anesthetics - mainly remifentanil and propofol infusions. One of these studies described a case of possible malignant hyperthermia (without triggering agents); however, the diagnosis of malignant hyperthermia in that case was questionable.

Our preference is to avoid triggering anesthetics and instead to use a total intravenous anesthesia (TIVA) technique.

There are no reports of using regional anesthesia in patients with MD-CMD. If a regional technique is chosen, careful consideration should be used due to the presence of spine deformities and contractures. The use of ultrasound could be helpful in these situations but has not been reported to date.
Necessary additional diagnostic procedures (preoperative)

Pulmonary:
If possible, pulmonary function tests. If not possible, determine child's needs for: cough assist, noninvasive ventilation support (i.e. CPAP), and mechanical ventilation.

If there is a possibility that assisted ventilation will be necessary postoperatively, CPAP should be started preoperatively. Also, evaluate type and functioning of tracheostomy.

Cardiac:
Pre-operative EKG, echocardiogram and cardiology consultation should always be done. Particular attention should be placed on assessing conduction abnormalities such as right bundle branch block in the EKG. In the echocardiogram, look specifically for left ventricular dysfunction (left ventricular dysfunction present in 1/3 of MD-CMD patients), dilated cardiomyopathy and signs of pulmonary hypertension.

Particular preparation for airway management

Difficult airway should be expected due to poor control of the head and marked hypotonia of neck and trunk muscles. Other contributing factors include: limited mouth opening due to jaw contractures, as well as maxillary discrepancy and macroglossia.

Particular preparation for transfusion or administration of blood products

Preparation for typical blood loss that occurs in neuromuscular spine fusion cases.

Particular preparation for anticoagulation

Not reported.

Particular precautions for positioning, transport or mobilisation

Patients with MD-CMD have multiple deformities of the limbs, joints and spine, including neck contractures, scoliosis, hip dislocation and subluxation. These require extreme precaution when transporting, mobilizing and positioning the patient. Extra-padding in pressure areas may be needed.

Probable interaction between anaesthetic agents and patient’s long term medication

Not reported.
Anaesthesiologic procedure

Pre-anesthetic evaluation:
In addition to the cardiovascular and respiratory evaluation described above, assess neurologic involvement (type, frequency and treatment of seizures) and gastrointestinal function (swallowing difficulties and risk of aspiration).

Anesthetic management:
- Prepare for difficult airway management (fiberoptic intubation is recommended)
- Tailor ventilatory support according to degree of respiratory compromise (most patients present with severe restrictive pulmonary disease). Anticipate needs for post-operative ventilatory support.
- Avoid use of depolarizing muscle relaxants due to potential hyperkalemic cardiac arrest and rhabdomyolysis. Use non-depolarizing muscle relaxants only if strictly necessary. Titrate non-depolarizing muscle relaxants and opiates carefully given baseline hypotonia.
- Controversy still exists around the use of halogenated agents. Our preference is to avoid these agents. However, exposure to halogenated agents might be necessary to place an intravenous catheter, in extremely challenging situations.
- Tailor anesthetic management according to degree of cardiovascular compromise. Severe cardiovascular instability could be expected depending on type of surgical procedure and cardiac function.

Particular or additional monitoring
In addition to standard ASA monitors, monitoring of neuromuscular blockade is highly recommended given the degree of baseline hypotonia. For spine cases with severe scoliosis, consider transesophageal echocardiogram (TEE) or central venous pressure (CVP) monitoring. Patients with MD-CMD are susceptible to cardiovascular instability due to poor left ventricular function and mechanical obstruction of the right ventricle in the prone position.

Possible complications
Possible anesthetic complications include:
- Cardiac arrest from hyperkalemia and rhabdomyolysis (secondary to the use of depolarizing muscle relaxants).
- Postoperative respiratory insufficiency requiring prolonged mechanical intubation
- There is not sufficient evidence to judge whether or not the risk of an MH-like reaction is increased, but given that MH-like reactions have occurred in myopathic patients in the absence of halogenated anesthetics and in the absence of succinylcholine all equipment needed to identify and treat MH should be available.
• Hypoglycemia.

• Cardiovascular instability should be expected even in asymptomatic patients (reports in the literature describe that at least half of patients with cardiac abnormalities do not report any symptoms).

Postoperative care

It is important to anticipate the possibility of respiratory problems postoperatively. Extubation to noninvasive ventilation (i.e. CPAP) should be considered, but one must always prepare for the possibility of respiratory failure and ICU admission. The degree of preoperative pulmonary and cardiac involvement will often dictate what type of postoperative care is necessary. Other contributing factors may include the patient’s nutritional status and degree of scoliosis.

Information about emergency-like situations / Differential diagnostics

causes by the illness to give a tool to distinguish between a side effect of the anaesthetic procedure and a manifestation of the diseases, e.g.:

There are no common emergency-like situations associated with this disease. However, one must be vigilant for signs of malignant hyperthermia given the possibility of its occurrence with or without triggering agents.

Ambulatory anaesthesia

Ambulatory anaesthesia is not recommended due to cardiovascular and respiratory compromise.

Obstetrical anaesthesia

There are no reports of obstetrical anesthesia care for these patients (probably due to the short life span secondary to disease complications).
Literature and internet-links


Last date of modification: August 2013

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