Anaesthesia recommendations for patients suffering from

Multiple pterygium syndrome, Escobar variant

| Disease name: | Multiple pterygium syndrome, Escobar variant, OMIM#265000 |
| ICD 10: | Q79.8 |
| Synonyms: | Escobar syndrome; pterygium syndrome; multiple pterygium syndrome, non-lethal type; familial pterygium syndrome; pterygium colli syndrome; pterygium universale |

Multiple pterygium syndrome, Escobar variant (MPSEV) is a rare congenital condition, which is inherited with an autosomal recessive pattern. It has an unknown incidence but is more common among children from consanguineous relationships. It is caused by a mutation in the CHRNG gene, on chromosome 2q. This gene encodes the gamma subunit of the acetylcholine receptor (AChR), which is found in the fetus until around 33 weeks gestation, when it is replaced by another subunit to form the adult AChR protein. The severity of the CHRNG gene mutation influences the severity of the condition and the complete absence of the gamma subunit will result in lethal multiple pterygium syndrome.

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

The clinical features of MPSEV are variable, but the condition is characterised by pterygia (excessive webbing) typically affecting the neck, axilla, digits (syndactyly), antecubital fossa, popliteal and intercrural areas. Akinesia, which may be identified before birth, also frequently results in arthrogryposis (congenital contractures), causing a crouched stance, and muscle weakness. Other typical findings include growth retardation, ankyloglossia (adhesions between the tongue and the palate), syngnathia (congenital bands of tissue between the maxilla and the mandible), cleft palate, lumbar lordosis and scoliosis.

Typical facial features seen in patients with MPSEV are ptosis, down slanting palpebral fissures, epicanthal folds, micrognathia, long philtrum and low set ears.

Other orthopaedic manifestations or associations that have been reported include cervical spine fusion, rib fusion, hip dislocation, foot deformities, camptodactyly, absent patellae, and pectus excavatum. There are case reports describing abnormalities of the ear bones and conductive hearing loss and two case reports describe patients with MPSEV who had cardiac defects (atrial septal defects). Males with this condition can have cryptorchidism and females can have missing or underdeveloped labia majora. People affected by MPSEV may also have respiratory distress at birth due to lung hypoplasia.

Unlike other abnormalities of the AChR, MPSEV does not lead to myaesthetic symptoms in later life as the gamma subunit is only expressed during fetal life. However, progressive webbing and scoliosis commonly cause significant reduction in lung capacity and an airway that is increasingly difficult to manage.

Eventhough myasthenic features and abnormal muscle histopathology is not expected in CHRNG mutations, however, congenital diaphragmatic muscle weakness, diffuse myopathy, and myasthenic-like features have been frequently reported in CHRNG mutation patients which could be due to the role of γ-subunit AChR in muscle organogenesis.

Typical surgery

- Orthopaedic surgery - scoliosis correction, tendon transfers, joint surgeries, contracture releases
- Plastic surgery - cleft palate repair, removal/reduction of pterygia, syndactyly release
- Ophthalmic surgery - surgery to improve ptosis
- Urological surgery - orchidopexy

Type of anaesthesia

The type of anaesthesia will depend on the severity of the symptoms and the surgical procedure. General, regional and local anaesthetic techniques are safe to use.

A case report does describe a case complicated by malignant hyperthermia but several other case reports have described the uneventful use of volatile anaesthetics. Kachko et al. presented a case using epidural anaesthesia and stated that the risk of malignant hyperthermia in MPSEV is low.

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There is little research on the function of the subunits of the adult AChR in MPSEV patients and some authors have avoided neuromuscular blocking agents due to concerns about their duration of action. However, case reports have described the uneventful use of both vecuronium and atracurium.

Short-acting agents, such as remifentanil, propofol and desflurane may be preferred in these patients as residual anaesthetic effects may impact postoperative upper airway control and respiratory function.

Where possible, regional or local anaesthesia should be used as an adjunct to general anaesthesia to reduce the use of drugs causing respiratory depression. Anatomical features such as kyphoscoliosis, lumbar lordosis and contractures may make regional anaesthesia and local anaesthetic blocks technically challenging and so the use of ultrasound should be considered to aid placement.

### Necessary additional diagnostic procedures (preoperative)

Prior to anaesthesia, patients with MPSEV should have a thorough preoperative assessment with particular attention paid to assessing the airway, and the respiratory, musculoskeletal and cardiovascular systems.

Additional preoperative investigations will be dependent on the severity of the underlying disease and the surgical procedure. Specific investigations to be considered include:

- Nasendoscopy to evaluate airway anatomy
- Arterial blood gases and lung function tests to assess the degree of pulmonary involvement
- Chest X-ray to help assess for cardiorespiratory disease and assess the extent of scoliosis
- Neck X-ray due to the risk of cervical spine stiffness and fusion
- ECG and echocardiogram to assess for underlying cardiac defects and disease

### Particular preparation for airway management

A difficult airway should be anticipated in all cases of MPSEV and so a comprehensive plan for a difficult airway should be in place prior to induction of anaesthesia. The airway should be assessed thoroughly pre-operatively including assessing mouth opening, micrognathia, ankyloglossia, neck webbing and neck movement.

The characteristics of MPSEV may make a standard facemask fit poorly and a 2 person technique may be required. The ability to accomplish adequate bag-valve-mask ventilation should be demonstrated prior to giving neuromuscular blocking agents.

Appropriate equipment for dealing with the difficult airway must be readily available and it may be necessary to have an ENT surgeon on standby to perform a tracheostomy as a last resort.

Unlike other congenital syndromes where management of the airway becomes easier, in MPSEV intubation becomes more difficult due to increased deformity of the airway by the pterygia. An airway that has previously been easy to manage can become increasingly
difficult with time. Detailed documentation of previous approaches to the airway is essential as these patients often present for multiple surgeries during their lifetime. Education about their airway should be provided to the patient and family in case they have to receive care at an outside or unfamiliar hospital.

**Particular preparation for transfusion or administration of blood products**

There is no definite recommendation for transfusion. Administration of blood products will depend on the type of surgery, the patient symptoms and advice from haematology clinicians.

**Particular preparation for anticoagulation**

There is no evidence to support the need for particular anticoagulation. Severe contractures may cause impaired mobility in patients and may suggest a higher risk of postoperative thrombosis.

**Particular precautions for positioning, transport or mobilization**

Caution must be taken while positioning the patient due to multiple contractures and possible scoliosis. Position should ideally be guided by the awake and cooperative patient.

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

None reported.

**Anaesthesiologic procedure**

The anaesthetic of patients with MPSEV should be managed by an experienced anaesthetic team, familiar with the management of difficult airways. A comprehensive airway plan should be made and appropriate equipment and personnel available in case of difficulty.

Both inhalational and intravenous induction have been used successfully. When inhalational induction is used, intravenous access should be established prior to induction. Intravenous access may be difficult due to anatomical abnormalities and intraosseous access or central venous access may be required.

Extubation should be done in an awake patient wherever possible.

**Particular or additional monitoring**

Routine monitoring is needed in all MPSEV patients. Flexion contractures may make it difficult to obtain intravenous access and non-invasive blood pressures and this may mean the use of invasive monitoring and central venous access is necessary. Monitoring of
neuromuscular blockade is recommended in all patients in whom neuromuscular blocking drugs are used.

**Possible complications**

Patients with multiple pterygium syndrome are at increased risk of failed or difficult intubation and ventilation may be difficult due to restrictive lung disease.

These patients may be sensitive to opiates and are of high risk of postoperative respiratory complications.

**Postoperative care**

Postoperative care will depend on the surgery performed and the severity of the clinical manifestations of the disease. Postoperative admission to an intensive care or high dependency care unit may be required for monitoring or ventilatory support, given the risk of postoperative respiratory failure. Mobilization can prove difficult due to contractures and postoperative opiates should be used with caution.

**Information about emergency-like situations / Differential diagnostics**

*caused by the illness to give a tool to distinguish between a side effect of the anaesthetic procedure and a manifestation of the disease*

All anaesthetists anaesthetising patients with Escobar should be familiar with difficult and failed intubation protocols.

**Ambulatory anaesthesia**

The suitability of ambulatory anaesthesia will be guided by the severity of disease, the surgical procedure and the local guidelines. It is unlikely to be appropriate in any but the mildly affected.

**Obstetric anaesthesia**

Pregnant women with MPSEV should have experienced consultant led obstetric and anaesthetic care and multidisciplinary team involvement.

In the only case report of the obstetric anaesthetic management of a lady with MPSEV, the patient underwent an elective Caesarean section under general anaesthesia. Neuraxial anaesthesia was not possible due to extensive lumbosacral surgery with placement of hardware. Due to a potentially difficult airway an awake fibreoptic intubation was performed and the anaesthetic was otherwise uneventful.

There are no case reports on labouring women with MPSEV but the likely problem of a difficult airway suggests that the early implementation of epidural anaesthesia would be beneficial. However, spinal abnormalities may make this technically very challenging or impossible.

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Literature and internet links

11. Online Mendelian Inheritance in Man (OMIM), http://www.ncbi.nlm.nih.gov/Omim/ (for Escobar or multiple pterygia syndrome, MG, congenital myasthenic syndrome, CHRNA1, CHRNB1, CHRND, CHRNE, and CHRNG)

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