Anesthesia in Mowat-Wilson disease: information on 11 Italian patients

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Abstract

Mowat-Wilson syndrome is a genetic disease caused by heterozygous mutations or deletions of the ZEB2 gene and characterized by typical clinical features. The congenital malformations typical of this syndrome call for early diagnostic and surgical procedures requiring general anesthesia, but few information about the anesthesiology management of such patients is available. We enrolled 11 families of patients with Mowat-Wilson syndrome who had undergone surgical or diagnostic procedures requiring general anesthesia, and sent them a retrospective questionnaire including 16 open questions about the procedures. They were further contacted by phone for a semi-structured interview. A total of 37 procedures requiring general anesthesia was reported in 11 patients. Only two patients reported anesthesia-related complications during the procedure. No true additional anesthesia-related risk was present for the patients with MW syndrome, besides difficult intubation, weaning and lower respiratory tract infection. Perception of risk, however, is derived by non-medical observation on the part of the parents.

Introduction

Mowat-Wilson syndrome (MWS; OMIM#235730) is a genetic disease caused by heterozygous mutations or deletions of the ZEB2 gene, and characterized by typical clinical features, including Hirschsprung disease (HSCR), agenesis of the corpus callosum (ACC), congenital heart disease (CHD), genital anomalies (particularly hypospadias in males), and eye defects. Since the first delineation by Mowat et al. (1998), more than 200 patients with ZEB2 mutations, deletions or cytogenetic abnormalities have been reported primarily from Northern Europe, Australia, Italy and the United States, and over 100 mutations have been described.3-17

Congenital malformations typical of this syndrome call for early diagnostic investigations and surgical resolutions; general anesthesia is required for diagnostic and surgical procedures, both in election and in emergency, but few information about anesthesiologic management is available.9

The aim of our study was to collect retrospective data on general anesthesia interviewing a large numbered group of Italian patients with MWS, assess the anesthesiologic outcomes and identify the conditions that required general anesthesia; aware of the fact that few clinical data would be derived from this interview.

Materials and Methods

A retrospective cohort study design was performed with questionnaire was sent to the parents using the mailing list of the Italian association of the Mowat-Wilson Syndrome. Thirty six questionnaires were sent out in total and each questionnaire included 16 open questions about age, type of surgery/diagnostic test requiring general anesthesia and outcome of the procedure. Some questions relating to the perception of the families about the knowledge of the disease by the anesthesiologist were also included. All of the families who initially replied to the questionnaire by mail were then contacted by phone for an additional semistructured interview.

Furthermore a further call was necessary to obtain clinical data such as the hospital stay from the clinical documentation in the possession of the families.

Results

We have collected questionnaires from 11 patients with MWS on thirty six.

A total of 37 anesthesiologic procedures was reported in 11 patients; in all cases, anesthesia was required in relation to various type of surgery and diagnostic procedures (Table 1).

The age of surgery and diagnostic procedures is reported in Figure 1.

The more common operations were the following: gastrointestinal surgeries (27%); diagnostic investigation, mainly MRI (24,3%); cardiac surgeries (16,2%); orchiopexy/orchiectomy of a cryptorchid testicle (13,5%); hypospadias correction (5,4%) (Figure 2). Unfortunately it was not possible to derive from the interviews the duration of the single surgical and instrumental procedures. 75.7% of families reported that the anesthesiologists were not aware of the syndrome; nevertheless, 43.2% of parents declared that they have been sufficiently reassured by the competence of the physician to trust him/her, even if we do not have data that attest to the clinical experience of anesthesiologists with pediatric patients.

In 78,4% of cases no complications of anesthesia emerged during the procedures (data collected from patients’ perception of patients and not from clinical data). Two patients reported anesthesia-related problems: a patient had cardiac complication during one procedure and another patient had respiratory complications during two
Anesthesia was required for 33 elective procedures and for 4 urgent procedures. No one of the parents received a report after the procedure.

### Discussion

The involvement of patients’ organizations in collecting clinical data has been a strong instrument for natural history of rare diseases. The anesthetic risk is present at each procedure but particular syndromes may provide an additional risk. However, very little is known about the anesthetic risk of MWS. The survey through social media has advantages and disadvantages, as well summarized in a recent article by William Davies: fast turnaround, low costs, a large number of patients recruited (often seen individually in different hospitals) but the evaluation of the point of view of the individuals or the caregivers, the compilation of retrospective questionnaire was on a voluntary basis and this may introduce an additional bias.

However, a similar investigation made for the MWS was done in the same manner for the Cri du Chat syndrome. Analysis of the type of surgery allows to note that most of these are made in election, except for those made in urgency (i.e. digestive canal at birth), when the diagnosis of MWS has not yet been made.

No true additional anestheisia-related risk was present for the patients with MW syndrome, both parents and professionals can take comfort in knowing the real anesthesiologic risk linked to issues involving the Mowat-Wilson syndrome. It is recommended to carry out an examination of patient’s cardiac function before any procedure; in case of cardiac malformation, antibiotic prophylaxis for bacterial endocarditis is required. Furthermore, in accordance with Deshmukh et al., the main anesthetic challenges in MW syndrome are a risk of difficult intubation and possibility of difficulty in weaning, lower respiratory tract infection and anemia.

The recollections of the patient’s parents may be biased, even if it was their choice whether or not to participate in the study, hence we can assume that their responses correspond to reality, but not as a real clinical evaluation.

### Table 1. Age of surgery and diagnostic procedures in 11 Italian patients with MWs.

<table>
<thead>
<tr>
<th>Patient</th>
<th>Age</th>
<th>Type of diagnostic/ surgical procedure</th>
<th>Elective/ urgent procedure</th>
<th>Patient’s trust in the anesthesiologist</th>
<th>Complications</th>
<th>Hospital stay</th>
</tr>
</thead>
<tbody>
<tr>
<td>PT1</td>
<td>3 yr</td>
<td>megacolon resection</td>
<td>elective</td>
<td>excellent</td>
<td>no</td>
<td>9 d</td>
</tr>
<tr>
<td>PT2</td>
<td>18 mo</td>
<td>brain MRI</td>
<td>elective</td>
<td>good</td>
<td>no</td>
<td>2 d</td>
</tr>
<tr>
<td>PT3</td>
<td>52 d</td>
<td>megacolon resection</td>
<td>urgent</td>
<td>very good</td>
<td>no</td>
<td>11 d</td>
</tr>
<tr>
<td>PT4</td>
<td>21 d</td>
<td>diagnostic bronchoscopy</td>
<td>elective</td>
<td>satisfactory</td>
<td>no</td>
<td>2 d</td>
</tr>
<tr>
<td>PT5</td>
<td>3 mo</td>
<td>cardiac catheterization + PDA closure</td>
<td>elective</td>
<td>very good</td>
<td>no</td>
<td>6 d</td>
</tr>
<tr>
<td>PT6</td>
<td>16 mo</td>
<td>adenotonsillectomy</td>
<td>elective</td>
<td>very good</td>
<td>no</td>
<td>5 d</td>
</tr>
<tr>
<td>PT7</td>
<td>1 mo</td>
<td>colostomy confection</td>
<td>elective</td>
<td>satisfactory</td>
<td>no</td>
<td>3 d</td>
</tr>
<tr>
<td>PT8</td>
<td>1 mo</td>
<td>pulmonic valve stenosis surgery</td>
<td>urgent</td>
<td>very poor</td>
<td>respiratory</td>
<td>5 d</td>
</tr>
<tr>
<td>PT9</td>
<td>1 mo</td>
<td>megacolon</td>
<td>urgent</td>
<td>good</td>
<td>no</td>
<td>7 d</td>
</tr>
<tr>
<td>PT10</td>
<td>6 mo</td>
<td>MRI</td>
<td>elective</td>
<td>satisfactory</td>
<td>no</td>
<td>2 d</td>
</tr>
<tr>
<td>PT11</td>
<td>11 mo</td>
<td>tear duct opening + orchidopexy</td>
<td>elective</td>
<td>satisfactory</td>
<td>no</td>
<td>5 d</td>
</tr>
</tbody>
</table>

**Note:** The table provides a detailed list of surgical procedures performed on 11 Italian patients with MWS, including the patient’s age, type of procedure, whether it was elective or urgent, the patient’s trust in the anesthesiologist, complications, and hospital stay. The table reflects the diversity of surgical procedures and the anesthetic considerations involved in managing these patients.
Conclusions

The study presents possible limits, such as potential bias due to the set nature of the design: a retrospective survey based on the memory of the patients’ parents. Moreover, the lack of some clinical data such as the duration of surgical and instrumental procedures, ASA score, the experience of the anesthetists inherent in pediatric patients, the anesthetic technique used, do not make the present study definitive. However our perception of risk, however, is derived by non-medical observation on the part of the parents, therefore further investigations – including the analysis of the medical files of all subject – are needed to confirm our findings; a prospective study would be desirable.

References