Abstract

Background and objectives: Malignant hyperthermia (MH) is a pharmacogenetic disease that causes abnormal hypermetabolic reaction to halogenated anesthetics and/or depolarizing muscle relaxants. In Brazil, there is a hotline telephone service for MH since 1991, available 24 hours a day in São Paulo. This article analyzes the activity of the Brazilian hotline service for MH in 2009.

Methods: Prospective analysis of all phone calls made to the Brazilian hotline service for MH from January to December 2009.

Results: Twenty-two phone calls were received: 21 from the South/Southeast region of Brazil and one from the North region. Fifteen calls were requests for general information about MH. Seven were about suspected MH acute episodes, two of which were not considered as MH. In five episodes compatible with MH, all patients received halogenated volatile anesthetics (2, isoflurane; 3, sevoflurane) and one also used succinylcholine; there were four men and one woman, with a mean age of 18 years (2-27). The problems described in the five MH episodes were tachycardia.
Introduction

Malignant hyperthermia (MH) is a pharmacogenetic disorder with autosomal dominant inheritance, characterized by abnormal hypermetabolic response to inhalational anesthetics (halogenated group) and depolarizing muscle relaxants, such as succinylcholine. Typical episodes feature tachycardia, tachypnea, hypercarbia, muscle rigidity, hyperthermia, metabolic acidosis, and rhabdomyolysis. However, there is great clinical variability, with abortive, atypical or fulminant forms, and even isolated masseter spasm. MH also has genetic variation: approximately 50% to 70% of cases are associated with ryanodine receptor gene mutations (a calcium release channel from the sarcoplasmic reticulum into the cytoplasm of the muscle fiber), but mutations have been described in other genes related to calcium metabolism, such as the dihydropyridine receptor gene. MH was first described in 1960 and occurs at frequencies up to 1:10,000 general anesthesias in children and 1:50,000 in adults. The first report on MH in Brazil occurred during halothane anesthesia and was published in 1975. However, there is no data on the actual incidence of MH in Brazil, a continental country with 190 million inhabitants. Considering the 3,890,755 anesthesias performed at the Unified Health System (SUS) in 2009 and 1:50,000 frequency of MH episodes during general anesthesia in adults, one would expect at least 77 reports of MH episodes during this period.

In 1991, a service for MH was created in Brazil called Hotline (+55-11-55759873), which is available 24 hours per day. The Brazilian hotline service for MH is located in the city of São Paulo at Hospital São Paulo, Escola Paulista de Medicina, Universidade Federal de São Paulo (EPM/Unifesp). Between 1991 and 2008, phone calls to the hotline were directed to the Division of Surgical Intensive Care and answered by the intensive care staff, assisted by MH supervisors. Since 2009, phone calls have been redirected to a group consisting of two supervisors - also specialized in MH investigation (HCAS, neurologist - and JLGA, anesthesiologist) - and eight doctors. The physicians received training that consisted of an intensive course on MH, with theoretical and practical information. Training included dantrolene dissolution and an intensive course on MH, with theoretical and practical information. Training included dantrolene dissolution and in vitro muscle contracture test (IVCT) in response to halothane and caffeine. Initial categorization and tabulation of data on call location, caller, description of call reason, proposed solution, consultant identification, and date. For classification, two independent investigators analyzed each record. In 2009, the hotline service for MH was later continued in two patients by in vitro muscle contracture test.

Conclusions: The number of calls per year in the Brazilian hotline service for MH is still low. The characteristics of MH episode were similar to those reported in other countries. The knowledge of MH in Brazil needs to be increased.

Material and methods

This was a prospective, analytical and observational study. The study’s primary material was the attendance records of the Brazilian hotline service for MH in 2009. These attendance records consisted of data identifying the caller, call location (city, state, hospital service, or residence), return phone number, description of call reason, proposed solution, consultant identification, and date. For classification, two independent investigators analyzed each record. Initially, categorization and tabulation of data on call location, caller, description of call reason, and solution proposed were performed. For records in which the call reason was a suspected episode of MH, we used a clinical scale described by Larach et al. to grade the likelihood of MH. This scale should not be used to make a diagnosis of MH susceptibility, but serves to classify the severity of episodes. In this 1- to 6-point scale (from almost incompatible to almost certain), MH suspected episodes are classified according to points obtained when analyzing the following variables: muscle rigidity, rhabdomyolysis, hyperthermia, cardiac abnormality, acid-base changes, and response to treatment with dantrolene. The study secondary material was the medical records of patients who, after being notified of the Brazilian hotline service for MH in 2009, were referred to Cedhima for evaluation/investigation and provided signed informed consent (CEP 0970/08). Data collected from medical records were demographic; medical, family, and epidemiologic history; current complaints; general physical examination; neurological examination; and laboratory tests, including histopathological study of muscle with histochemistry and in vitro muscle contracture test in response to halothane and caffeine. Histopathological study of muscle included the following reactions and staining: hematoxylin-eosin, periodic acid-Schiff (PAS), modified Gomori trichrome, Sudan black B; succino dehydrogenase (SDH), cytochrome C-oxidase (COX), nicotinamide adenine dinucleotide tetrazolium reductase (NADH), and adenosine triphosphatase (ATPase) acidic (pH 4.3) and alkaline (pH 9.4), according to the methodology described by Dubowitz.

In vitro muscle contracture test response to halothane and caffeine was performed according to the protocol of the European Malignant Hyperthermia Group (www.emhg.org). After biopsy, four fragments (0.5x2 cm) of vastus lateralis muscle are immediately placed into a Krebs-Ringer’s solution (95% oxygen and 5% carbon dioxide) and dissected into thin fragments (2-3 mm) under stereomicroscope (Olympus, USA). Then, each fragment is attached to a silver electrode at one
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end and to a force transducer at the other (transducer Narco Bio Systems, International Biomedical, USA). Tests are made in baths heated to 37°C, through thermostatically controlled circulating water (Mecânica Castro Saray, Brazil), filled with Krebs-Ringer’s solution and continuously carboxygenated. Each fragment is continuously stimulated with supramaximal electrical pulses of 1 ms and frequency of 0.2 Hz (Boeck stimulator). The degree of muscle contraction is monitored in real time during the entire test and detected by force transducer linked to an IBM 486 computer, using an amplifier/analog-digital converter and program for biological data acquisition Aquados (Links, Brazil). At least four tests are performed for each patient: two with caffeine and two with halothane. To test caffeine, concentrations of 0.5, 1.0, 1.5, 2.0, 3.0, 4.0 and 32.0 mmol.L⁻¹ are used. To test halothane, the concentrations are 0.5, 1.0, 2.0 and 3.0%. Test results are referred to as the threshold, which is the lowest concentration of caffeine and halothane producing a minimum increase of 0.2 g in basal tension.

According to test results, patients are classified into three groups:

a) **Malignant hyperthermia susceptibility (MHS):** caffeine threshold concentration of 2.0 mmol or less and halothane threshold of 2.0% or less;

b) **Malignant hyperthermia nonsusceptible (MHN):** caffeine threshold concentration of 3.0 mmol or more and halothane threshold of 3.0% or more;

c) **Malignant hyperthermia equivocal (MHE):** other results are considered equivocal.

From a practical standpoint, MHE patients are also considered susceptible to MH.

**Results**

In 2009, the Brazilian hotline service for MH received 22 calls: 21 from the South/Southeast region and one from the North region (Pará State). The Southeast region had 17 calls, all from the State of São Paulo, and the Southern region had four calls, three from Paraná and one from Rio Grande do Sul.

Among the 22 calls, 15 were requests for general information about MH and seven about suspected MH acute episodes. General information consisted on how to purchase dantrolene (4), how to research MH susceptibility (3), and how to obtain additional information about the disease (8). For the general information, proposed solutions were, respectively: contact the customer service department (SAC) of the lab selling dantrolene in Brazil (Cristália: +55-11-0800 7011 918); seek MH outpatient clinic to schedule a visit (telephone +55-11-55719667); and visit Cedhima’s website (MH section) at www.anestesiologia.unifesp.br. The additional information requested by the eight callers were: how to assemble a hospital kit of MH; who may have MH; which is the safe anesthesia for MH; where to find literature on MH for school work; and, finally, if halothane exhaled by a pet dog after surgery could be dangerous for susceptible individuals to MH. The first three information are available on Cedhima’s MH section, while the fourth information initially recommended keeping the animal in the veterinary clinic during postoperative period and, subsequently, a consultation with MHAUS (Malignant Hyperthermia Association of the United States: www.mhaus.org.br) was done and it was suggested to follow the recommendation of the MHAUS consensus about the lack of risk, to susceptible individuals, associated with residual anesthetic in the operating room air, provided there is the necessary exhaustion. Table 1 shows the distribution of calls by caller’s characteristic.

Among the seven patients with suspected MH acute episode, three calls came from the hospital that houses the Brazilian hotline service for MH; afterwards, two of these patients underwent investigation and susceptibility to MH was confirmed. Given that 16,865 anesthetic procedures were performed at Hospital São Paulo (EPM/Unifesp) in 2009 (including anesthetics for surgeries and diagnostic procedures), the presence of two MH episodes would indicate, in this service, a frequency of 1.1:10,000. Of these, 8,906 were general anesthesia, which would imply a MH episode frequency of 2.2 per 10,000 general anesthesia cases in this service.

Considering the seven patients with suspected MH episode after reviewing the history and additional tests required, in two of them the episodes were not considered as MH. One patient had not used triggering agents (halogenated or succinylcholine), and the event was later diagnosed as a reaction to fentanyl. The other patient had postoperative hyperthermia and sepsis was diagnosed.

All five patients whose episode was considered compatible with MH underwent anesthesia with halogenated anesthetics (2 with isoflurane, 3 with sevoflurane); one patient also received succinylcholine. This group consisted of four men and one woman, mean age 18 years (range 2-27). Surgeries performed were orthopedic (1), ENT (ears, nose, throat) surgery (1), gastrointestinal (2), and thoracic (1). The problems described in these five patients during the MH episode were, in order of frequency: tachycardia (5 patients: mean of 130 bpm, range of 120-140 bpm); increase in expired carbon dioxide (4 patients: mean of 70 mm Hg, range of 52-96 mm Hg); hyperthermia (3 patients: mean of 39.5°C, range of 39-40°C); acidemia (1 patient: pH 7.22); rhabdomyolysis (2 patients: CPK 4,701-27,037 U.L⁻¹); and choluria (1 patient). According to the clinical grading scale to assess the likelihood of MH, these five MH episodes had a mean score of 28 points (range 18-43). Table 2 presents the clinical grading scale to assess the likelihood of MH and anesthetics used in five patients.

Relatives of two of the patients considered compatible with MH reported that deaths had occurred during anesthesia in other family members.

<table>
<thead>
<tr>
<th>Table 1 Characteristics of Callers.</th>
<th>Number (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Anesthesiologist</td>
<td>8 (36%)</td>
</tr>
<tr>
<td>Intensivist</td>
<td>1 (4.5%)</td>
</tr>
<tr>
<td>Orthopedist</td>
<td>1 (4.5%)</td>
</tr>
<tr>
<td>Physician</td>
<td>2 (9%)</td>
</tr>
<tr>
<td>(specialty not specified)</td>
<td></td>
</tr>
<tr>
<td>Nurse</td>
<td>1 (4.5%)</td>
</tr>
<tr>
<td>Pharmacist</td>
<td>4 (18%)</td>
</tr>
<tr>
<td>Graduate student (medicine)</td>
<td>1 (4.5%)</td>
</tr>
<tr>
<td>Patient</td>
<td>3 (14%)</td>
</tr>
<tr>
<td>Ignored</td>
<td>1 (4.5%)</td>
</tr>
<tr>
<td>Total</td>
<td>22</td>
</tr>
</tbody>
</table>
Guidance to MH compatible individuals was done according to the informed characteristic of the episode and standardized treatment protocol. In four patients, MH episode ceased with discontinuation of triggering halogenated agents and measures such as cooling (2 patients) and hyperventilation with 100% oxygen (2 patients); in one patient (number 3) dantrolene was administered. All five patients were monitored in the intensive care unit and recovered without sequelae; two of them were referred for evaluation and investigation at Cedhima (patients 2 and 4) and will be described in detail below.

Patient 2: male, aged 27 years, ASA I, with history of tumor mass in the left thigh, whose biopsy diagnosed alveolar sarcoma. The patient received four cycles of chemotherapy (ifosfamide and doxorubicin), which have been completed 10 months ago, and subsequently amputation was administered. All five patients were monitored in the intensive care unit and recovered without sequelae; two of them were referred for evaluation and investigation at Cedhima (patients 2 and 4) and will be described in detail below.

Subsequently, a new pulmonary metastasectomy was indicated (now to the left), with balanced/combined general anesthesia induced with midazolam, fentanyl, propofol, and atracurium. Isoflurane (up to 1 MAC) in addition to epidural anesthesia were used for maintenance. After induction, the patient evolved to persistent and unexplained sinus tachycardia, normothermia, and maximum expired CO2 of 52 mm Hg (Figure 1). Postoperatively, the patient developed diffuse myalgia and choloria. Serum creatine phosphokinase (CPK) levels were measured on the third postoperative day and showed a value of 27,037 U.L⁻¹ (normal value 190 U.L⁻¹), which raised the hypothesis of postoperative rhabdomyolysis due to MH. The patient evolved without further complications; was discharged from the ICU on the sixth day, informed about his possible susceptibility to MH, and referred to Cedhima’s outpatient clinic.

On physical and neurological examination during outpatient evaluation, the patient presented diminished breathing sounds in the left hemithorax and left lower limb amputation without sequelae. Two tests with halothane showed contracture (0.24 g and 0.48 g, concentration of 2%), which is consistent with positive results, indicating susceptibility to MH. Histopathological examination of skeletal muscle showed mild subsarcolemmal mitochondrial proliferation. Since the MH episode, the patient had already undergone two lung metastasectomies under total intravenous anesthesia, uneventfully.

Patient 4: female, aged 23 years, ASA I, with history of left patella chronic dislocation since 11 years old after fall from own height. The patient had undergone three previous uneventful surgical procedures (varus osteotomy, femorodistal pseudo-arthrodesis, and medial patellofemoral reconstruction), all on the left side. In the preanesthetic evaluation, the patient denied significant family history.

Subsequently, the patient was admitted for revision of left femur osteosynthesis evaluation, and general anesthesia was chosen, given the time scheduled for the procedure. Concomitant epidural anesthesia was initially attempted; however, due to accidental dural puncture, this procedure was abandoned. General anesthesia was induced with sufentanil, propofol, and pancuronium, with isoflurane and sufentanil used for maintenance. After four hours of uneventful surgery, without apparent surgical reasons, the patient showed increased capnometric values (up to 66 mm Hg, confirmed with arterial blood gas, also showing respiratory acidosis); tachycardia (121 bpm); and

<table>
<thead>
<tr>
<th>Score</th>
<th>Theoretical probability</th>
<th>Theoretical risk of MH</th>
<th>Patients and score (n)</th>
<th>Triggering anesthetic</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>Almost impossible</td>
<td>1</td>
<td>0</td>
<td></td>
</tr>
<tr>
<td>3 to 9</td>
<td>Unlikely</td>
<td>2</td>
<td>Patient 1: 18</td>
<td>Sevoflurane</td>
</tr>
<tr>
<td>10 to 19</td>
<td>Somewhat less than likely</td>
<td>3</td>
<td>Patient 2: 18</td>
<td>Isoflurane</td>
</tr>
<tr>
<td>20 to 34</td>
<td>Somewhat more than likely</td>
<td>4</td>
<td>Patient 3: 28</td>
<td>Sevoflurane + succinylcholine</td>
</tr>
<tr>
<td>35 to 49</td>
<td>Quite likely</td>
<td>5</td>
<td>Patient 4: 33</td>
<td>Isoflurane</td>
</tr>
<tr>
<td>50 or +</td>
<td>Almost certain</td>
<td>6</td>
<td>Patient 5: 43</td>
<td>Sevoflurane</td>
</tr>
</tbody>
</table>

Figure 1  Patient 2: ETCO2, Temperature, and Heart Rate during Intraoperative Period.
temperature elevation (baseline: 36°C, maximum reached 40°C, intraesophageal temperature sensor). The diagnosis of malignant hyperthermia was then hypothesized by the anesthesiologist, and the proposed measures were taken. The surgical team was notified of the suspected diagnosis and the maintenance anesthesia was replaced by exclusively intravenous injection of propofol via an infusion pump and opioid fentanyl, when necessary. Additionally, infusion of cold saline and hyperventilation with oxygen 100% was started, with improvement within minutes following these measures. After surgery and uneventful extubation in the operating room, the patient was transferred to the ICU for 48 hours. CPK showed progressive elevation from the surgery, with maximum of 4,701 U.L⁻¹, two days after surgery, but with gradual return to baseline. The patient was discharged on the sixth day after the procedure, asymptomatic, informed about the suspected susceptibility to MH, and referred to Cedhima’s outpatient clinic.

During outpatient evaluation, the patient reported a history of exertion intolerance, with cramps, in addition to congenital clubfoot. She reported several changes in her family: son with inguinal hernia; father with pectus carinatum, clubfoot, and chronic patellar dislocation; paternal aunt and grandmother with chronic patellar dislocation; first-degree cousin who died during surgery. On physical and neurological examination, the patient presented short neck, bilateral ptosis, and arched palate. Serum CPK was increased at rest (1,061 U.L⁻¹). Three months after discharge, she underwent right vastus lateralis muscle biopsy for in vitro muscle contracture test in response to halothane and caffeine. There was contracture in two tests with halothane (3.6 g and 1.56 g, concentration of 2%; Figure 2) and two tests with caffeine (2.92 g and 1.2 g, concentration of 2 mmol; Figure 3) compatible with positive results, indicating susceptibility to MH. Pathological examination of skeletal muscle showed sparse fibers with various changes, such as necrosis, hypertrophy, centralization or segmentation.

Discussion

Calls to MH hotline were mostly from the Southeast Region and within the State of São Paulo. This finding may be due to the fact that the hotline service is located in São Paulo and because there is specific legislation for MH in this state, which requires notification of the MH episodes and dantrolene availability. On the other hand, the complete lack of calls from the states of Rio de Janeiro and Santa Catarina could be explained by the centralization of services related to MH in these states (Universidade Federal do Rio de Janeiro and the Universidade Federal de Santa Catarina, respectively). However, it is more difficult to explain the lack of calls from the Northeast and Midwest regions, from Minas Gerais and Espirito Santo States, as well as a single call from the Northern region. With the increased level of suspicion, it is possible that more MH diagnoses will be made, as occurred in the hotline service for MH. In the past, MH was believed as being exclusive to Caucasians, which has proven to be incorrect with the description of MH episode in BLACKS, Orientals, and Indians. In Brazil, the initial impression that the MH genetic trait would be concentrated in European descendants in the south has also been disproved by genetic studies, such as the description of a patient from Minas Gerais who was compound heterozygous (i.e., an individual with two different mutations in their ryanodine receptor, each inherited from one of the non-consanguineous parents). This finding from the work of Kossugue et al. conforms the estimated frequency of the ryanodine gene mutation in ryanodine carriers, which could reach up to 1:2000. This prevalence would be much higher than the estimated rate for MH episode - explained by the fact that many mutation carriers are never anesthetized with triggering drugs and, even if exposed, they may not develop a MH episode, as occurred in patient 2 of this report.
Much of the calls were requests for information, which is understandable due to MH rarity, with characteristics of an orphan disease. Orphan diseases are rare and affect a small number of individuals and/or have no treatment that stimulates the interest of industry investments, diagnosis, and treatment.

Regarding dantrolene issues, in Brazil there is a Resolution of the Federal Council of Medicine (Resolution CFM 1.802/2006), which includes dantrolene on the list of “drugs that should necessarily be available in any setting where anesthesia is used” 22. Considering an individual weighing 70 kg and a loading dose of dantrolene 2.5 mg·kg–1, nine vials of dantrolene 20 mg would be required for the first bolus. Because bolus is repeated until the MH episode is controlled and up to 10 mg·kg–1 (4 boluses) may be needed to control it, 36 vials of dantrolene should be kept in stock. This stock must be immediately available to anesthesiologists, as the time to dilute dantrolene can reach 4 minutes per vial and the mortality rate is demonstrated to increase when the first dose is administered 30 minutes after the onset of the episode 22. Furthermore, there is a need for options for quickly acquiring additional dantrolene, in case there is a need for more than 10 mg·kg–1 of dantrolene to control a episode, a need to continue treatment for 24-48 hours, and the increasingly common possibility of overweight patients.

MH differential diagnosis includes various conditions that may lead to hypercapnia (problems in the ventilation circuit), hypermetabolism (pheochromocytoma, thyrotoxic episode, exogenous intoxication with salicylates), and/or hyperthermia (iatrogenic heating, sepsis). Although MH may occur postoperatively, it is unlikely that fever alone characterizes MH 25. Another cause for excluding MH diagnosis in hotline calls in 2009 was an isolated reaction characterized by muscle rigidity after opioid use, a rare event associated with high doses, which occurs during induction or recovery and may show improvement after naloxone or neuromuscular block 26.

Regarding MH episode presentation in our country, there are similarities with those reported in other countries. The halogenated agents involved were sevoflurane and isofluorane. Keep in mind that all halogenated agents may cause MH, even when the patient has not received succinylcholine 27. There was a clear predominance of male (80%), as well as young patients, similar to the recent revision by Larach et al. 24. Considering a patient weighs 70 kg and loading dose of dantrolene 2.5 mg·kg–1, nine vials of dantrolene 20 mg would be required for the first bolus. Because bolus is repeated until the MH episode is controlled and up to 10 mg·kg–1 (4 boluses) may be needed to control it, 36 vials of dantrolene should be kept in stock. This stock must be immediately available to anesthesiologists, as the time to dilute dantrolene can reach 4 minutes per vial and the mortality rate is demonstrated to increase when the first dose is administered 30 minutes after the onset of the episode 22. Furthermore, there is a need for options for quickly acquiring additional dantrolene, in case there is a need for more than 10 mg·kg–1 of dantrolene to control an episode, a need to continue treatment for 24-48 hours, and the increasingly common possibility of overweight patients.

The non-use of dantrolene in some reports may be associated with difficulty in obtaining the drug in some locations or instant improvement of abortive’s episode with discontinuation of halogenated agents. However, patients who do not receive dantrolene, even without presenting with cardiopulmonary arrest, are at risk for complications, such as liver failure or myalgia and rhabdomyolysis, as the case with patient 2 of this report 24.

Regarding clinical features associated with susceptibility to MH, several authors have reported osteoarticular changes (chronic dislocation, clubfoot), dimorphisms (cleft palate, ptosis, strabismus), and myopathies (central core myopathy). In this study, patient 4 illustrates this association, with a history of multiple dimorphisms and presence of osteoarticular changes over four generations 1,7,17.

Conclusions
Considering Brazil as a whole, the number of calls to the hotline service for MH is still small for a one-year period and there is a large concentration in the Southeast region. Moreover, the frequency of calls was proportionally much higher in the hospital housing the hotline service, probably due to the continuing education programs developed there. Awareness of MH in Brazil needs to be increased, in addition to actions already implemented (regular insertion of MH theme in anesthesiology meetings; production of updated material, such as books and Brazilian guidelines for MH, and development of electronic site for MH: http://www.unifesp.br/dcir/anestesia/hipertermiamaligna/index.html).

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References