



# Malignant hyperthermia and the clinical significance of type-1 ryanodine receptor gene (*RYR1*) variants: proceedings of the 2013 MHAUS Scientific Conference

## Hyperthermie maligne et signification clinique des variants du gène du récepteur de la ryanodine de type 1 (*RYR1*): comptes rendus de la conférence scientifique 2013 de la MHAUS

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**Abstract** *The Malignant Hyperthermia Association of the United States and the Department of Anesthesia at the University of Toronto sponsored a Scientific Conference on November 1-2, 2013 in Toronto, ON, Canada. The multidisciplinary group of experts, including clinicians, geneticists, and physiologists involved in research related to malignant hyperthermia (MH), shared new insights into the pathophysiology of diseases linked to the type-1 ryanodine receptor gene (RYR1) as well as the*

*relationship between MH and “awake MH” conditions, such as exertional rhabdomyolysis and exertional heat illness. In addition, the molecular genetics of MH and clinical issues related to the diagnosis and management of disorders linked to RYR1 were presented. The conference also honoured Dr. David H. MacLennan for his contributions to our understanding of the genetics, pathogenesis, and treatment of MH and other RYR1-related myopathies. This report represents a summary of the proceedings of this conference.*

**Author contributions** *Sheila Riazi, Sheila Muldoon, and Robert Dirksen were organizing committee members. Henry Rosenberg was Chair of the organizing committee. Sheila Riazi, Sheila Muldoon, Robert Dirksen, and Henry Rosenberg participated in article design. Sheila Riazi, Sheila Muldoon, Robert Dirksen, Henry Rosenberg, Natalia Kraeva, James Dowling, Clara Ho, Maria-Alexandra Petre, and Jerome Parness participated in manuscript preparation. Sheila Riazi is the archival author. Natalia Kraeva, Clara Ho, and Maria-Alexandra Petre participated in taking notes during presentations.*

**Résumé** *L'association américaine sur l'hyperthermie maligne (MHAUS: Malignant Hyperthermia Association of the United States) et le Département d'anesthésie de l'université de Toronto ont parrainé une conférence scientifique les 1<sup>er</sup> et 2 novembre 2013 à Toronto (ON, Canada). Le groupe multidisciplinaire d'experts, incluant*

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*des cliniciens, des généticiens et des physiologistes impliqués dans la recherche sur l'hyperthermie maligne (HM), a partagé de nouvelles connaissances sur la physiopathologie de la maladie liée au gène du récepteur de la ryanodine de type 1 (RYR1) ainsi que sur les rapports entre la HM et la « HM éveillé », telle que la rhabdomyolyse secondaire à un effort physique et la maladie de chaleur à l'effort. De plus, la génétique moléculaire de l'hyperthermie maligne et les problèmes cliniques liés au diagnostic et à la gestion des troubles associés au gène RYR1 ont été présentés au cours de la conférence. Le docteur David H. MacLennan y a été également honoré pour ses contributions à notre compréhension de la génétique, de la pathogénie et du traitement de l'hyperthermie maligne et des autres myopathies liées au gène RYR1. Ce compte rendu présente un résumé des travaux de cette conférence.*

Malignant hyperthermia (MH) is an autosomal dominant pharmacogenetic disorder triggered by exposure to volatile anesthetics and/or depolarizing muscle relaxants.<sup>1,2</sup> Malignant hyperthermia manifests as a potentially lethal hypermetabolic crisis associated with a rapid and uncontrolled increase in myoplasmic  $\text{Ca}^{2+}$  in skeletal muscle cells.<sup>3</sup> Since MH was first recognized as an inherited condition in 1960,<sup>4</sup> six potential genetic loci for MH susceptibility have been reported,<sup>5</sup> but specific MH-associated mutations have been identified in only two genes thus far, type-1 ryanodine receptor gene (*RYR1*; OMIM® 180901)<sup>A</sup> encoding the  $\text{Ca}^{2+}$  release channel of the sarcoplasmic reticulum (SR) and *CACNA1S* (OMIM 114208) encoding alpha-1S subunit of the voltage-dependent L-type calcium channel of the transverse tubule. Products of these genes play key roles in the process of excitation-contraction (EC) coupling and in the maintenance of  $\text{Ca}^{2+}$  homeostasis in skeletal muscle cells. Malignant hyperthermia genetic research has shown that *RYR1* is the primary causal gene for MH; *RYR1* mutations are found in 60-86% of MH families with diverse ethnicity.<sup>6,7</sup>

Genetic testing for MH plays an important role in MH diagnostics today. Genetic testing has proven to be especially useful in the early diagnosis of children and patients who will not undergo the caffeine-halothane contracture test (CHCT) for MH susceptibility because of the invasive nature of the muscle biopsy that the CHCT requires. Nevertheless, the genetic testing exhibits low sensitivity due to the limited number of mutations that have fulfilled the criteria for MH causation as determined by the

European Malignant Hyperthermia Group (EMHG; [www.emhg.org](http://www.emhg.org)) and the Malignant Hyperthermia Association of the USA (MHAUS; [www.mhaus.org](http://www.mhaus.org)).

In an effort to advance our understanding of MH and other *RYR1*-related diseases and to promote new insights into MH pathophysiology, diagnosis, and treatment, MHAUS, along with the University of Toronto, Department of Anesthesia, co-sponsored the MHAUS Scientific Conference held on November 1-2, 2013 in Toronto, Canada. The multi-disciplinary group of experts, including clinicians, geneticists, and physiologists involved in research related to MH, shared new insights into the pathophysiology of diseases linked to *RYR1*, the relationship between MH and “awake MH” conditions (e.g., exertional rhabdomyolysis [ER] and exertional heat illness [EHI]), the molecular genetics of MH, as well as other clinical issues related to the diagnosis and management of *RYR1*-linked disorders. A strong emphasis was on the clinical implications of recent advances. The conference honoured Dr. David H. MacLennan for his contributions to our understanding of the genetics, pathogenesis, and treatment of MH and other *RYR1*-related myopathies.<sup>7</sup>

The objectives of the 2013 MHAUS Scientific Conference were to share the latest research on:

- Type-1 ryanodine receptor (RyR1) structure/function and assessment of *RYR1*-related myopathies
- Molecular genetic diagnosis of MH
- Relationship of MH with EHI, ER, and statin myopathies
- Role of rare diseases and genetic variation databases
- Improved education in MH diagnosis and treatment

The key concepts and theories presented at the conference as well as the trends and future challenges in the field of MH research and patient care are summarized in Tables 1 and 2.

### **Malignant hyperthermia, *RYR1*-related myopathies and related conditions**

Structural and functional studies of the ryanodine receptor

Dr. F. Van Petegem (University of British Columbia) presented novel structural mechanisms by which disease-related *RYR1* mutations disrupt RyR1 function. Analysis of pseudo-atomic models and high-resolution structures of N-terminal domains of the ryanodine receptor indicate that disease mutations weaken interdomain interactions in RyR1, thus lowering the energetic barrier to channel opening and leading to enhanced calcium release.<sup>8</sup> Such studies, currently limited by the inability to crystallize the

<sup>A</sup> Online Mendelian Inheritance in Man. Available from URL: <http://omim.org/> (accessed August 2014).

**Table 1** Key concepts and theories presented at the scientific conference

- Combining cryoEM and *x-ray* crystal structure analyses can model effects of N-terminal disease mutations on RyR1 function and mechanisms of *RYR1*-related diseases, thereby clarifying mechanisms by which disease-causing mutations act to alter RyR1 function.
- Parallels can be drawn between mechanisms by which *RYR1* and *RYR2* disease mutations alter excitation-contraction coupling that lead to diseases in skeletal muscle (MH) and cardiac muscle (CPVT).
- Increasingly, the *RYR1* gene mutations are implicated in various inherited myopathies, statin-induced myopathy, and heat/exercise-induced rhabdomyolysis.
- Clarification is necessary regarding which *RYR1*-related myopathies predispose to MH.
- Identification of genes involved in MH, other than *RYR1* and *CACNA1S*, is required before comprehensive genetic testing for MH is possible.
- The metabolic basis of MH needs to be better understood.
- More work needs to be done to clarify which *RYR1*-related myopathies predispose to MH.
- The diagnostic algorithm for MH was reviewed with a focus on the characteristics of the clinical presentation, IVCT/CHCT, and genetic testing.
- MH therapeutic options are expanding, including a new formulation of dantrolene and the possible utility of newer drugs such as AICAR and carvedilol derivatives.
- A diagnostic algorithm for myopathic patients was presented.
- The continued evolution of genetic databases will be instrumental in advancing improved understanding of the genotype/phenotype relation in MH.
- Next-generation sequencing has the potential to identify new MH-causative genes and to improve the sensitivity and efficiency of MH genetic testing, and it will be instrumental in a more accurate assessment of the prevalence of MH susceptibility within the general population.

AICAR = 5-aminoimidazole-4-carboxamide ribonucleoside; CPVT = catecholaminergic polymorphic ventricular tachycardia; cryoEM = cryoelectron microscopy; IVCT/CHCT = *in vitro* contracture test/caffeine-halothane contracture test; MH = malignant hyperthermia; *RYR1* = type-1 ryanodine receptor gene

entire RyR1 protein, will aid in the design of new drugs for the treatment of MH and other *RYR1*-related disorders.

Dr. C. Franzini-Armstrong (University of Pennsylvania), a well-known expert on skeletal muscle ultrastructure, presented a study elucidating a role of the type-3 ryanodine receptor (RyR3). RyR3 is expressed in neonatal skeletal muscles and can also be found at low levels in a variety of cell types, but not adult mammalian skeletal muscle. Nevertheless, the physiological role of RyR3 in non-mammalian muscle remains unclear. Dr. Franzini-Armstrong, working on zebrafish skeletal muscle, showed that RyR3 forms parajunctional structures, “feet”, adjacent to the triad junction that serve to enhance  $Ca^{2+}$  sparks initiated by RyR1.<sup>9</sup>

**Table 2** Research directions and patient care issues

- Further characterization of RyR1 structure will likely elucidate pathophysiological mechanisms for MH and other *RYR1*-related disorders.
- New alternative treatments for acute MH and related crises are under investigation (AICAR, carvedilol derivatives, altered formulations for dantrolene).
- Improved understanding of the pathologic potential of *RYR1* variants will be facilitated by next-generation sequencing and new genetic databases such as ClinVar.
- The need to identify the metabolic underpinnings of MH should be facilitated by a metabolomics approach.
- Access to rare diseases databases promoted by collaborations with the Office of Rare Diseases Research and their Global Rare Diseases Patient Registry and Data Repository may reveal new links between MH and other rare disease entities.
- The need exists for further elucidation of the relationship between exercise-induced rhabdomyolysis, heat stroke, MH, and *RYR1* variants.
- The role of dantrolene and other MH therapies in ER and EHI should be explored.
- The importance of preoperative identification of MH risk among patients suffering from congenital myopathies should be advocated.
- Areas for further study include the epidemiology of MH across the globe and increased preparedness for treatment of MH in other countries and non-hospital sites where anesthesia is administered.

AICAR = 5-aminoimidazole-4-carboxamide ribonucleoside; ER = exertional rhabdomyolysis; EHI = exertional heat illness; MH = malignant hyperthermia; *RYR1* = type-1 ryanodine receptor gene

### Molecular mechanisms of RyR-related disorders

Excitation-contraction coupling is the process whereby an electrical signal in the T-tubule membrane (an action potential) is converted into an intracellular chemical signal (a  $Ca^{2+}$  transient) that is used to drive muscle contraction. Excitation-contraction coupling is composed of a series of steps in which a conformational change in the voltage-dependent L-type calcium channel induced by depolarization triggers activation of the RyR1  $Ca^{2+}$  release channel in the adjacent terminal cisternae of the SR. While the effects of MH mutations in RyR1 on EC coupling are well established, alterations in several other calcium signalling mechanisms have also been implicated in MH. In a process known as store overload-induced calcium release (SOICR), RyR1 spontaneously releases  $Ca^{2+}$  from the SR once luminal  $Ca^{2+}$  concentrations reach a critical threshold. Recent evidence suggests that RyR1 mutations may lower the threshold for SOICR, thus promoting spontaneous SR release of  $Ca^{2+}$ . Human Embryonic Kidney 293 (HEK293) cells expressing RyR1 receptors bearing the porcine MH mutation, R615C, displayed a reduced threshold for SOICR in comparison with wild type RyR1, a response that was potentiated by halothane and inhibited by dantrolene.<sup>10</sup> Important

parallels can be drawn between the RyR1-mediated mechanisms in MH and RyR2-mediated mechanisms in catecholaminergic polymorphic ventricular tachycardia (CPVT), a stress-triggered condition leading to bidirectional ventricular tachycardia and sudden cardiac death.

The keynote speaker of the conference, Dr. W. Chen (University of Calgary), discussed similarities between MH and CPVT. Dr. Chen's studies suggest that both disorders are caused by RyR mutations that reduce the threshold for SOICR at baseline, which is then further depressed by their respective triggers to cause severe spillover of  $\text{Ca}^{2+}$  into the cytoplasm, ultimately leading to uncontrolled and sustained contraction of the myofibrils.<sup>11</sup>

A recent study<sup>12</sup> identified yet another mechanism of RyR1 regulation by  $\text{Ca}_v1.1$ , the pore-forming subunit of the L-type  $\text{Ca}^{2+}$  channel encoded by *CACNA1S*.  $\text{Ca}_v1.1$  functions not only to activate RyR1 during EC coupling but also to suppress resting RyR1-mediated  $\text{Ca}^{2+}$  leak from the SR. Disruption of this  $\text{Ca}_v1.1$  inhibitory regulation of RyR1  $\text{Ca}^{2+}$  leak represents a novel molecular mechanism for increased sensitization of muscle cells to MH triggers.

In addition to mediating calcium release from the SR, RyR1 is also implicated in sarcolemmal calcium entry processes. One such  $\text{Ca}^{2+}$  entry process is known as excitation-coupled calcium entry in which calcium enters the cell through  $\text{Ca}_v1.1$  channels in the T-tubule membrane in response to membrane depolarization. Excitation-coupled calcium entry is not only significantly enhanced in myotubes expressing MH mutants of RyR1 but is also suppressed to wild type levels following exposure to dantrolene.<sup>13,14</sup> A second mechanism of  $\text{Ca}^{2+}$  entry, store-operated calcium entry (SOCE), recruits extracellular  $\text{Ca}^{2+}$  entry when intra-sarcoplasmic stores become depleted, as occurs during sustained contractions observed during an MH crisis. Myotubes from MH mouse models exhibit an increased rate of SOCE channel activation.<sup>15</sup>

Dr. J. Parness (Children's Hospital of Pittsburgh) focused on the question of potential explanations for incomplete penetrance in the presentation of MH, i.e., occurring after several uneventful exposures to volatile anesthetics. Two mechanisms were proposed: 1) involvement of a prion-like structure in a volatile-anesthetic-sensitive metabolic response element; 2) an inducible metabolic genetic element that is normally silenced but capable of being activated by MH-triggering agents and/or elevated  $\text{Ca}^{2+}$ . Dr. Parness suggested that engagement of metabolomic approaches, through detection of uniquely elevated metabolites during an MH episode, could reveal MH target proteins, genes, and metabolic pathway(s) involved in MH hypermetabolism.

## Malignant hyperthermia and other RYR1-related myopathies

Dr. J. Dowling (University of Toronto) brought attention to the expanding spectrum of *RYR1*-related myopathies that, in addition to MH, include congenital core myopathies (central core disease and multimini-core myopathy) and a growing collection of recessive non-core myopathies. Analysis of 106 cases with recessive *RYR1* mutations revealed their histologic heterogeneity - from classic core myopathies to non-core myopathies.<sup>16</sup> In pediatric populations, *RYR1* mutations are associated with extremity and axial weakness and often include ophthalmoplegia and facial weakness. Unlike dominantly inherited MH and central core disease (CCD), where mutations are often located in "hot spots" of *RYR1*, recessive mutations are spread throughout the *RYR1* gene. Cases with severe clinical phenotype were significantly associated with hypomorphic *RYR1* mutations, which result in decreased RyR1 protein expression.<sup>16</sup> It was suggested that increasing expression of the RyR1 protein might ameliorate the severity of the disease in such cases. The risk for MH in patients with those mutations is uncertain.<sup>17</sup>

Dr. Dowling discussed two recently identified causes of non-*RYR1* core myopathies. One was caused by a mutation in *MYH7* (a gene encoding myosin heavy chain beta isoform)<sup>18</sup> and another by a splice site mutation in *CCDC78* (a gene encoding the coiled-coil domain-containing 78 protein that plays a role in skeletal muscle contraction) validated in a zebrafish model. Dr. Dowling also presented a study that identified a recessive mutation in *STAC3* (a gene encoding the SH3 and cysteine-rich domain 3 protein) encoding a newly identified component of the EC-coupling machinery,<sup>19</sup> in patients with Native American myopathy, a rare myopathy associated with dysmorphic features and MH susceptibility. Based on studies on *RYR* zebrafish as well as on myotubes from patients with *RYR1* mutations, Dr. Dowling showed that antioxidant treatment reduces oxidative stress and improves motor function. Thus, the use of antioxidants may represent a viable therapeutic strategy for patients with *RYR1*-related myopathies.<sup>20</sup>

## Evaluation and anesthetic management of patients with myopathy

To minimize the risk of an adverse reaction in patients with myopathies, a preoperative patient evaluation should be conducted ideally in conjunction with a neurologist. Anesthesiologists, aware of an increased risk for MH in patients with *RYR1*-related congenital myopathies, should consider MH susceptibility in myopathic patients who

develop fever, tachycardia, hypercapnia, and/or hyperkalemia perioperatively very high on their differential diagnostic list and manage accordingly.

The presence of skeletal muscle weakness is a major perioperative concern, especially if it involves the respiratory and/or bulbar muscles. Although congenital myopathies are not usually associated with primary myocardial involvement, scoliosis-related restrictive lung disease can lead to heart failure. A formal clinical evaluation of respiratory and bulbar muscle function and cardiac reserve is a mandatory aspect of the preoperative evaluation of patients with congenital myopathies. There should also be a plan for available postoperative ventilatory support and a suitable postoperative intensive care unit.

Dr. M. Tarnopolsky (McMaster University) recommended evaluation of baseline levels of serum potassium and creatine kinase (CK) and levels of aspartate aminotransferase and alanine transaminase as key indicators of a propensity for profound perioperative rhabdomyolysis.<sup>21</sup> Interpretation of CK levels requires consideration of the variability of normal CK values, with CK of 800 U·L<sup>-1</sup> being the upper limit for African Americans but only 200 U·L<sup>-1</sup> being the upper limit for Caucasians.<sup>22</sup> Dr. Tarnopolsky outlined a diagnostic algorithm for evaluating patients with myopathies and suggested that patients presenting with an abnormal neurological exam (e.g., unexplained chronic high CK, any CK over 1,000, positive family history of high CK), neuromuscular disorder, or non-hypertensive cardiomyopathy, should be sent for further testing (i.e., muscle magnetic resonance imaging and/or muscle biopsy as well as genetic testing). He also pointed out that special considerations are warranted for patients with myotonic muscular dystrophy type 1 with cardiac conduction defects and slower rates of drug metabolism, for patients with Duchenne, and for those with other severe forms of muscular dystrophy who may have cardiomyopathy and may be particularly susceptible to electrolyte disturbances such as hyperkalemia and sudden cardiac death.

One important discussion point was whether a non-triggering anesthetic, i.e., total intravenous anesthesia, should automatically be administered to myopathic children. Participants agreed that patients with multiminicore or central core disease should avoid triggering anesthetics, children and adults with significant muscle weakness due to primary myopathies should have their functional status assessed before the operation, and transfer to specialty centres with intensive care facilities should be considered. While all agreed that succinylcholine should be avoided in such patients, a blanket prohibition of volatile anesthetics in any and all myopathies was not considered appropriate. In conclusion, a fully informed decision concerning the use of volatile anesthetics in a patient with myopathy includes a thorough analysis of family and personal history and requires

an understanding of the underlying molecular defect, as different underlying pathophysiological mechanisms would have different implications for anesthetic management of patients and their families.

### Relationship of MH with EHI, ER, and statin myopathies

The link to heat/exercise-induced rhabdomyolysis

Similar to *RYR1*-related myopathies, EHI and ER may share a common pathogenic mechanism with MH susceptibility. The relationship between EHI/ER and MH was explored in studies presented by Drs. F. O'Connor, J. Capacchione, and N. Sambughin of the Uniformed Services University of the Health Sciences (USUHS). One military study showed that 15 out of 26 unrelated active-duty males diagnosed with recurrent or unexplained ER had a positive CHCT, and that nine of 13 who underwent genetic screening carried an *RYR1* variant.<sup>23</sup> Dr. P. Hopkins (Leeds Malignant Hyperthermia Unit, UK) found that many patients recovering from heat stroke or ER and whose symptoms responded to dantrolene carried pathogenic *RYR1* mutations and tested positive on the halothane-caffeine *in vitro* contracture test (IVCT). Dr. F. O'Connor described studies from Dr. P. Deuster's lab at USUHS showing that some patients tolerated heat stress less than others and had a more prominent rise in post-exercise CK. Questions emerging from these studies include: 1) What percentage of individuals suffering from EHI/ER will exhibit pathologic *RYR1* variants? 2) What precautions should be taken with regard to heat/exercise in individuals susceptible to MH? 3) What is the potential utility of dantrolene in the treatment of EHI/ER?

A recent case of "awake MH" reported by Dr. Capacchione supported a link between EHI and MH.<sup>24</sup> An otherwise healthy six-year-old boy developed lower extremity rigidity, trismus, and fever after playing in a splash pool. On arrival in the emergency department, he appeared to be seizing. Use of succinylcholine led to cardiac arrest and death. Postmortem genetic analysis revealed a novel *RYR1* variant that was later found in his father who tested positive by CHCT and was diagnosed histologically with CCD.<sup>24</sup> Similar cases of "awake MH" in children, with and without recognized myopathies, have been described previously.<sup>25,26</sup>

Health care providers, especially first responders who use succinylcholine (e.g., emergency department physicians), should be aware of the risks of succinylcholine administration in patients, particularly children, presenting with muscle rigidity and hyperthermia, even in the absence of exposure to anesthetic drugs.

The observation that many individuals with EHI/ER are positive in either the CHCT or IVCT indicates that a subset of EHI/ER might represent a variant clinical expression of MH. It is not clear, however, whether individuals with documented episodes of MH are more susceptible to EHI. There are yet no biochemical, genetic, or functional tests to diagnose susceptibility to EHI/ER or to differentiate clearly between MH and EHI/ER. Unexplained, persistently elevated serum CK levels in the absence of neuromuscular disease should alert anesthesiologists and sports medicine physicians to a potentially greater than normal risk of MH and EHI/ER.<sup>23-28</sup>

#### The link between MH and statin myopathies

Dr. G. Vladutiu (University of Buffalo) discussed statin-induced myopathies (SIM). She pointed out that, although serious adverse myopathic reactions to statin therapy are rare, about 200,000 people in the U.S. (i.e., 0.5% of 40 million who are taking statins) are at risk for life-disabling myopathy, and 4-6 million (10-15%) develop muscle pain/weakness. In some cases, these symptoms become chronic.<sup>29,30</sup> A study by Dr. Vladutiu's team revealed that predisposition to SIM is a genetically complex condition associated with mutations in at least eight muscle disease genes, including *RYR1* and *CACNA1S*. Carrier status for certain autosomal recessively inherited myopathies is increased as much as 20-fold in cases of severe statin myopathy.<sup>31</sup>

Dr. S. Hamilton (Baylor College of Medicine) showed that some statins, such as simvastatin, interfere with intramyofibre  $\text{Ca}^{2+}$  homeostasis. Simvastatin triggered hypermetabolic responses in mice carrying an MH-associated Y524S mutation in *RyR1*, and this response was prevented by pretreatment with 5-aminoimidazole-4-carboxamide ribonucleoside (AICAR), possibly through its direct effect on *RyR1*.<sup>32</sup>

The potential role of statins as a risk factor for increasing the likelihood of an MH reaction during anesthesia in a malignant hyperthermia susceptible (MHS) individual remains unknown.

#### Clinical diagnosis of MH

The clinical presentation of MH is highly variable. Common early signs include hypercarbia (end-tidal  $\text{CO}_2 > 60$  mmHg), sinus tachycardia, masseter muscle rigidity, and generalized muscle rigidity. Hyperthermia was an early sign in only 8.2% of cases but was the first to third sign in 63.5% of cases.<sup>33,34</sup> Most commonly, MH presents intraoperatively within an hour of anesthetic induction, but it may also present in the early postoperative period and may recrudescence following treatment. Other signs of

MH include hyperkalemia-related changes in an electrocardiogram, ventricular ectopy or fibrillation, bigeminy, tachycardia, myoglobinuria, and disseminated intravascular coagulation (DIC). The likelihood of development of MH complications increases 1.6-fold for every 30-min increase in time period from the first sign of MH to the first dose of dantrolene and 2.9-fold for every  $2^\circ\text{C}$  increase in maximum temperature (T-max).<sup>33</sup>

Dr. M. Larach (North American Malignant Hyperthermia Registry) presented data on MH deaths collected in the North American Registry (NAMHR) from 2007-2012 and compared the results with the analysis of similar cases reported during 1987-2006.<sup>25</sup> Examination of the NAMHR for the earlier time period revealed 291 episodes of MH with eight cardiac arrests (2.7%), including four fatalities (1.4%). In analyzing 84 cases with seven cardiac arrests and eight deaths (9.5%), cases from the AMRA (adverse metabolic/musculoskeletal reaction to anesthesia) reports filed with the Registry from 2007-2012, important differences were found between MH death and survival groups: 1) The likelihood of death increases 14-fold with the use of a skin temperature probe or absence of any temperature monitoring and increases 4.2-fold with every  $1^\circ\text{C}$  increase in T-max, underscoring the importance of accurate real-time temperature monitoring during anesthesia. 2) Every 1 mEq rise of potassium increases the likelihood of death 4.5-fold. 3) Disseminated intravascular coagulation greatly increases the likelihood of death (odds ratio, 101.5). Compared with the earlier NAMHR study, there was an increased failure to rescue from MH even though these patients were, for the most part, young, healthy, and undergoing low- to intermediate-risk surgery. In an effort to improve our understanding of this potentially lethal condition, health care providers are encouraged to obtain blood specimens appropriate for molecular genetic analysis from all patients suffering a possible MH-related arrest or death. Whenever possible, muscle biopsy contracture studies should be performed on survivors of cardiac arrest that is potentially associated with MH.<sup>6,25</sup>

#### Update on caffeine-halothane contracture testing

The caffeine-halothane contracture test, known as CHCT in North America<sup>35</sup> and as IVCT<sup>36</sup> in Europe, remains the gold standard for MH diagnosis, being at present the most sensitive diagnostic test for MH susceptibility. As reported by Dr. S. Muldoon (Uniformed Services University), 60-80 CHCT tests are performed annually in five MH testing centres in North America using a standardized protocol.<sup>B</sup> Anesthesiologists at these centres review the medical,

<sup>B</sup> Malignant Hyperthermia Association of the United States. Available from URL: [www.mhaus.org](http://www.mhaus.org) (accessed August 2014).

**Table 3** Classification of variants obtained from NGS data using *RYR1* as an example\*

Variant	Definition	Example	Significance	Report
Pathogenic	Confirmed as a disease-causing mutation	R614C	MHS Diagnosis	Yes
Benign polymorphism	Natural variation in DNA with no known adverse effect	Variant observed in healthy populations with high frequency	Could be a genetic modifier in future studies	No
VUS	Low frequency variant of unknown significance	Missense variant in a less evolutionary conserved <i>RYR1</i> region	May change status to pathogenic with additional data	Yes
Incidental finding	Variant that carries a potential risk of a disease state but is not relevant to the patient's primary diagnosis	Carrier state for mutations in <i>CFTR</i> gene	Not diagnostic for MH. Assessment of clinically actionable defined by ACMG guidelines	Yes, if clinically actionable

\* Modified with permission from<sup>40</sup>

ACMG = the American College of Medical Genetics and Genomics, *CFTR* = cystic fibrosis transmembrane conductance regulator gene, MHS = malignant hyperthermia susceptibility, NGS = next generation sequencing, *RYR1* = type-1 ryanodine receptor gene, VUS = variant of unknown significance

family, and anesthetic history of each referral to determine the patient's prior probability of having an MH event. A Clinical Grading Scale (CGS) generated by a consensus of MH experts often aids in evaluating the likelihood of a patient having experienced an MH episode.<sup>37</sup> The CGS scores rigidity, muscle breakdown, respiratory acidosis, temperature increases, cardiac involvement, and family history to evaluate the likelihood of MH. The CGS requires careful documentation of the clinical events, temperature monitoring, and blood gas analysis in order to develop an accurate ranking.

The disadvantages of both contracture tests, i.e., their reliance on invasive muscle biopsies and the cost of maintaining testing centres, has led to the closure of several centres in the Western Hemisphere and to a decrease in the number of CHCT tests performed annually.

Dr. S. Riazi (University of Toronto) discussed the disparities of phenotype and genotype concordance results from the Canadian database. These studies found that 22 patients (from eight families, containing two probands) were identified as MHS by CHCT, yet they did not carry their familial *RYR1* variants during workup. Seventeen of these patients (from three families) lacked familial mutations identified as causative in the EMHG database. The preponderance of discordance can be explained either as false positivity of CHCT or a lower than expected reliability of the current genetic testing. Thus, caution should be taken in counselling family members based solely on genetic results, which emphasizes a) the complementary role of CHCT or IVCT analyses in making a definitive MH diagnosis and b) the need for more definitive identification of genes and physiology that

both confer MH susceptibility and define the hypermetabolic crisis that is MH.

#### *Advances in MH genetic testing*

A prominent focus of the conference centered on a discussion regarding how novel genetic analytical techniques can be used to provide insights into the complex genetic basis of MH. Next-generation sequencing (NGS) platforms enable fast and cost-efficient sequencing of all protein coding regions - whole exome sequencing (WES) - in the human genome. Whole exome sequencing based on NGS holds great promise for overcoming significant challenges in MH genetic research, such as MH genetic heterogeneity as well as the large and complex structures of the two known MH candidate genes, *RYR1* and *CACNA1S*.

A key challenge in using WES for patient diagnosis is the ability to distinguish between disease-causing variants and benign polymorphisms. The American College of Medical Genetics and Genomics has developed recommendations for interpretation of sequence variations using standardized terminology.<sup>38,39</sup> Genetic variants are typically grouped into four categories: pathogenic, benign, variant of unknown significance (VUS), and incidental findings<sup>40</sup> (Table 3).

Two studies presented at the conference revealed high levels of allelic heterogeneity in the *RYR1* and *CACNA1S* genes. Drs. Hopkins and Kim<sup>41</sup> analyzed *RYR1* and *CACNA1S* variants identified using WES in 5,379 individuals as part of the National Heart, Lung, and Blood Institute Grand Opportunity Exome Sequencing Project.

Their analyses showed that *RYR1* is more genetically diverse than 74–79% of other genes in European-Americans and African Americans, while *CACNA1S* is more genetically diverse than 77–82% of other genes in the two cohorts.<sup>41</sup> Additionally, the data analyses indicated that most of the rare *RYR1* variants detected - with frequency less than 1% - are likely to be benign polymorphisms. The study also showed that current *in silico* bioinformatics prediction tools discriminate poorly between pathological and presumed non-pathological *RYR1* variants.

In the study by Gonsalves *et al.*,<sup>42</sup> the authors evaluated exome sequencing data from the ClinSeq<sup>®</sup> study on a cohort of 870 volunteers not selected for MH susceptibility. The study identified 70 *RYR1* variants and 53 *CACNA1S* variants in this cohort. Remarkably, a known MH-causal mutation was detected in one study participant without a medical or family history of MH; the finding was reported to the patient's family doctor as an incidental finding. As in the previously described study,<sup>41</sup> most of the novel variants were predicted to be benign. The authors also reported that a significant proportion of *RYR1* sequence variants in the human gene mutation database (HGMD) were erroneously classified as “disease-causing mutations” due to the absence of adequate genetic (e.g., co-segregation) or functional data and should be reclassified as being benign, probably benign, or a VUS.

Further studies into creating useful algorithms to predict a given variant's pathogenicity are clearly warranted and would require additional clinical and genetic data such as segregation patterns in large families and validation of altered RyR1 function in either cell systems or genetically engineered animals. As of today, only 31 out of more than 300 *RYR1* coding variants are officially classified as MH-causative and, thus, are used for clinical genetic testing for MH.<sup>C</sup>

Dr. N. Sambughin (USUHS) reported the use of WES for genetic analysis of MHS patients negative for *RYR1* variants. Their study revealed that such patients represent a clinically and genetically heterogeneous group of muscle disorders that may carry variants in genes previously not implicated in MH susceptibility.

All the presenters in this session emphasized that, at present, targeted sequencing (TS) through a gene panel that uses the NGS platform seems to be the most practical approach for screening MH candidate genes, including *RYR1* and *CACNA1S*. Targeted sequencing has a higher coverage (up to 99%) and accuracy and, therefore, higher sensitivity than WES. Depending on the laboratory, TS panels are also supplemented with traditional Sanger

sequencing to “backfill” exons that are poorly or incompletely covered by NGS. This gives an added level of coverage and reduces the potential for false negative results. Thus, accurate MH diagnosis could be improved through the use of TS of *RYR1* and *CACNA1S* as part of an integrated approach that also includes careful consideration of the patient's anesthetic and family history, CHCT/IVCT results, genotype-phenotype correlations, histopathology, and clinical evaluation for neuromuscular disorders.

#### Update on treatment of MH

Dantrolene remains the only specific drug treatment for acute MH. Use of dantrolene along with greater awareness and education by such groups as MHAUS has led to a reduction of MH mortality from 80% in the 1960s/1970s to less than 10% today. A multicentre study presented by Drs. Pawar and Rosenberg showed that dantrolene is also effective in treating related MH-like syndromes, such as neuroleptic malignant syndrome (NMS), sepsis, and cocaine intoxication.<sup>43,44</sup> Dantrolene is a relatively safe medication, as indicated by an analysis of 368 AMRA reports submitted to the NAMHR.

Other potential drugs were also discussed at the conference. Dr. Hamilton reported that AICAR, an AMP-activated protein kinase activator, prevented heat-induced sudden death and decreased cytoplasmic Ca<sup>2+</sup> levels in a mutant RyR1 knock-in mouse model of MH.<sup>45</sup>

Additionally, given similar roles of RyR1 and RyR2 in skeletal and cardiac muscle, respectively, it is possible that carvedilol, which suppresses the store overload-induced Ca<sup>2+</sup> release in cardiac myocytes and prevents ventricular tachyarrhythmias in heart failure, may also be useful as a potential MH pharmacotherapy.<sup>10</sup>

Dr. J. Philips (West Virginia University) presented a poster emphasizing the importance of intraoperative cognitive aids to treat MH successfully. This field will likely be facilitated by the recent introduction of the MHApp. This iPhone application not only guides users through the steps of MH management but also generates a report of the event that can be filed with the NAMHR.

#### Integration of MH data with global databases of rare diseases

Malignant hyperthermia is one of approximately 6,000 rare diseases that together affect 6–8% of the global population.<sup>46</sup> Single medical research centres do not typically have access to a sufficient patient population on their own to conduct comprehensive clinical or translational research studies for a particular rare disease, hampering the development and validation of effective treatments for these conditions.

<sup>C</sup> European Malignant Hyperthermia Group. Available from URL: <https://emhg.org/genetics/> (accessed August 2014).



In 2010, the Office of Rare Diseases Research (ORDR) at the National Institutes of Health initiated the creation of a Global Rare Diseases Patient Registry (GRDR) and the development of a repository of rare disease biospecimens.<sup>47</sup> Drs. B. Brandom (NAMHR) and D. Maglott (National Center for Biotechnology Information) discussed the role of the ORDR and genetic databases in MH risk assessment and as a resource for clinicians and scientists. Dr. Brandom reported that the NAMHR of MHAUS was selected to be one of 12 existing organizations to contribute data to the GRDR, a two-year pilot project with the aim of facilitating data analyses across different rare diseases and promoting the development of clinical trials and translational research for rare diseases.

Dr. Maglott emphasized the importance of genetic variation databases for clinical genetics. Next-generation sequencing has resulted in an exponential growth in the identification of genetic variants, making it difficult to keep up with new information added to genetic variation databases such as LOVD, OMIM, HGMD, and UniProt/Swiss-Prot. The newly released ClinVar database provides a comprehensive resource for disease variants (<https://www.ncbi.nlm.nih.gov/pubmed/24234437>).

The combination of the complex genetic nature of MH and the genetic diversity of *RYR1* and *CACNA1S* genes make these databases and bioinformatics tools particularly valuable in the assessment of the pathogenic potential of newly identified genetic variants.

In his presentation, Dr. H. Rosenberg, President of MHAUS, emphasized that the mission of MHAUS is to promote education, training, preparedness, and research related to MH and MH-like syndromes. The MH Hotline provides invaluable direct real-time assistance to clinicians dealing with potential MH cases. About 600 calls per year are fielded by the 32 volunteer hotline consultants. Although support for clinical and laboratory research through MHAUS funding is limited, the organization serves a critical role in sharing new information and insights among the clinical and patient communities, promoting initiatives to enhance MH preparedness and patient care, and reducing morbidity from MH. As a result of conferences such as this one supported by both MHAUS and the University of Toronto, Department of Anesthesia, the significance of *RYR1* variants in medicine beyond those related to anesthesia is better appreciated and, thus, will stimulate further advances that will benefit patients and clinicians alike. Malignant hyperthermia is probably the most dramatic example of how *RYR1* variants may lead to alteration of muscle physiology and biochemistry, but it is certainly not the only syndrome or disease state affected by this channel. The conference displayed a widespread interest in a better understanding of the role and

significance of RyR1 and related proteins in muscle physiology and a wide range of clinical syndromes.

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