Malignant Hyperthermia: The Spontaneous development of the MH-PSS Syndrome

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ABSTRACT

Mating MH positive pigs to MH positive pigs produces an F1 generation that is highly stress susceptible. Recording the MUP shows MH + pigs have a higher u voltage than control pigs. Older MH+ pigs have an even higher u voltage than control pigs. The duration of the voltage spike is also increased in MH+ pigs versus control pigs and older MH+ pigs have even a longer duration of the voltage spike.

We can assume that by concentrating the MH genetic defect in the F1 generation that the population of defective sodium channels in the acetylcholine receptor was present at a high concentration. Since the acetylcholine receptors are spatially located under the footpiece of the myoneural junction which makes them a bank of receptors that are readily accessible when acetylcholine is released by the nerve, and the action of acetylcholine is very rapid. Therefore, the electromyographic data reflects the genetically defective sodium channels as the major functional component when we recorded the data.

The sodium channels can be likened to a low voltage switch in a telephone circuit that is used to route telephone calls.

We would suggest that the sodium channel at the acetyl choline receptor has been adapted to produce heat as well as muscle contraction and that the ability to produce copious amounts of heat is the biological mechanism that differentiates warm blooded animals from cold blooded animals.

Introduction

The lethality of MH is directly related to the flow of Na+ ions thru the sodium channels in sufficient quantity to destroy the -90mv charge at the cell membrane, thereby overwhelming the cells Na+, K+ ATPase capacity to pump Na+ out of the muscle cell which subsequendy results in muscle cell death. The death of the muscle cells from the rapid production of large quantities of heat and acids denatures the cell proteins which thereby become nonfunctional. The acids and heat are picked up by the circulatory system, distributed to other parts of the body, and result in further damage to organs and other tissues which leads to the death of the whole organism.

We have been fortunate in being able to record full blown MH-PSS syndrome responses under controlled and recordable...
conditions. We are presenting those results in this manuscript. They are published in BMJ [4]. They are reprinted here for ease of comparison. With permission of BMJ.

These results indicate that the metabolic rate increased to over 10x normal with an intense peripheral vasoconstriction that shuts down heat loss via radiation which returns when the pig dies and the hot body is radiating heat at a high level.

In addition, we are presenting narrative accounts of the MH-PSS syndrome under field conditions.

The simplest way to increase MUP voltage is to lengthen the time the sodium channel is open. That is exactly what has happened in the adult prigs from our breeding colony. There may also be an increased flow of sodium ions thru the sodium channel when it is open.

Narrative Accounts

During the collection of calorimeter data in the Missouri Partitional Calorimeter we transported susceptible pigs from Sinclair Farm to Building T-13 in wood transport crates. The distance of travel was about 5 miles.

We loaded the first pig into the calorimeter and started recording data. In a few minutes, the pig developed MH_PSS and subsequently died in the calorimeter. The experimental results are in BMJ [3].

The key data are reprinted here to illustrate rapid onset and rampant acceleration of the metabolic rate and heat output. While the calorimeter was running, I went back out to the holding area to check on the other pigs which were awaiting their turn for the calorimeter run.

A white pig was basically in difficulty as I could hear the labored breathing and see the agitation experienced by the animal. I gently let the pig out of the transport crate. The pig circled around on the concrete floor 2 or 3 times then lay down and died. The white skin color had a purple mottled appearance that indicated the effect of an intense peripheral vasoconstriction.

Another pig, a large breeding boar died on a hot (105°F) day in the swine barn facility. All of these animals were posted by Loren Kintner, DVM who diagnosed the deaths due to MH-PSS.
Methods and Materials

The pigs used in these experiments were raised from the initial five pigs reserved as breeding stock at Arlington Farms, UW during January 1969. They were challenged with 3% Halothane in O2 and identified as MH susceptible whenever the rear legs became extended and appeared to be in rigor. The Halothane challenge was discontinued and the animals spontaneously recovered and were used for breeding stock. The two males and three females were raised in Swine Barn A under Leo’s supervision. Unfortunately, when they reached 225-240 lbs. in weight, they were hauled off to the UW slaughter house. Leo found them missing and calling the slaughter house and recovered one male and two females that had not been slaughtered. We moved the MH breeding animals to the UW Sheep Farm so there could not be any other possible confusion with other pigs in Swine Barn A. I moved the key MHS breeding animals to Sinclair Farm at U of MO in 1973 when I relocated to the University of Missouri for a new research position. Later in 1982, the entire stock of MH susceptible breeding animals was moved to TTUHSC at El Paso, TX. They were located on a pig farm near to El Paso, TX.

We found the Halothane challenge test to be very useful in identifying MH susceptible pigs for breeding and experimental work. By mating MHS + males with MHS+ females we have produced MHS++ F1 pigs that were highly susceptible to the MH-PSS genetic syndrome. The defective sodium channels at the myoneural junction are present at a high population level. The MH genetic defect is not sex linked so the reverse mating program produces highly susceptible pigs. Therefore, the electromyographic data reflects the genetically defective sodium channels as the major functional component when we recorded the data.

We believe that MH-PSS susceptibility is directly related to a higher MUP voltage and duration in the older animals. Our previous publication’s data is presented in graphical format for easier evaluation and comparison.

Duration of the voltage spike represents the length of time the sodium channel is open to allow sodium ions to pass through and create the voltage spike. The data suggests that the age of the animals plays a role in the duration of the spike. This observation is consistent with actual susceptibility to the MH-PSS syndrome whenever pigs are stressed by hauling to market. The Organon 9426 data indicated a reduction in the actual number of functional sodium channels in MHS versus normal pigs. Thus, aging makes the pigs more susceptible to develop the MH-PSS syndrome.

James R Stewart, an ATT Special Services telephone repairman [5], likens the sodium channel to a switch in a low voltage circuit that is used to route telephone calls. The sodium channels are spatially located under the foot piece of the motor nerve where they can act as bank of switches to regulate muscle function.

Further, we would suggest that the sodium channel futile cycle is most likely the biological mechanism used by warm blooded animals to maintain core temperatures at the appropriate level of 37-38°F. Basal metabolic rates may not be sufficient to maintain an elevated core body temperature. The prime example of that result would be cold blooded animals that must seek a warm environment in order to raise body temperature up to a functional level.

Our ability to control the mating in the pig colony allowed us to routinely produce homozygous F1 offspring. This provided us with very susceptible animals that exhibited the MH syndrome at maximal sensitivity.

Please note that the older MH susceptible pigs continued to exhibit an increased voltage spike and a longer duration before the sodium channels finally closed. We believe this increased voltage for a longer duration is the manifestation of genetically defective sodium channels. A 30% decrease in the actual number of sodium channels may be the result of a biological correction to enable the animal to survive with the MH syndrome genetic defect [2,6-9].
References