Extending the Fitzhugh Nagumo Excitable Tissue Model to Model Myocardial Necrosis

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Abstract

Neuroscience frequently relies on robust mathematical models to simulate and investigate trends in neuromechanics and signal propagation. We propose an examination of the Fitzhugh-Nagumo model, and additionally propose a modified extension of this model to account for necrosis of cardiac cells. We use this model to investigate case studies in cardiac necrosis and examine the effect of necrosis on a one dimensional cardiac model.

1 Examination of the Fitzhugh Nagumo Model

The modelling of neurons and associated action potentials started in the early 1900s with the "integrate and fire" model. In 1952, Hodgkin and Huxley created a biologically relevant 4 dimensional system of equations to describe neuron dynamics at a point. While very interesting to biologists, mathematicians found the model too difficult to systematically study. In 1961 Richard Fitzhugh created a 2D caricature of the Hodgkin Huxley model (activation and recovery) using a modified Van-der-Pol oscillator. This model has been both well studied and well characterized. Additionally, it has found many uses in cardiac models due to its simplicity and similarity in shape to a cardiac action potential.

The Hodgkin Huxley equations described how neuronal membrane voltage changed over time, given a large number of parameters.

$$C_m \frac{dV}{dt} = I_m = I_{Na} + I_K + I_l \tag{1}$$

$$I_x = \bar{G}_x p_{open}(V_m - E_x); x \in \{Na, K, l\}$$
(2)

$$p_{open_K} = n^4 (3)$$

$$p_{open_{Na}} = m^3 h (4)$$

$$p_{open_l} \equiv 1 \tag{5}$$

$$\frac{dx}{dt} = \alpha_x(V)(1-x) - \beta_x(V)x; x \in m, n, h$$
 (6)

(7)

This is impossible to solve analytically, and quite difficult to explore numerically due to the high dimensionality of the system and the large number of parameters. In studying hyperplanes through the 4D phase space, Nagumo noticed that all planes of activation and recovery variables had a similar shape, so he tried modeling neuronal activity as a modified Vander-Pol oscillator (Bonheffer-Van-der-Pol oscillator). This model, while not looking like stereotyped action potentials, has several features of the full hodgkin huxley model such as activation hysteresis and spike blocking. Despite not looking like a neuronal action potential, it closely resembles cardiac action potentials and have since been adapted as one of the de-facto models for cardiac simulations.

1.1 Model Rationale and Variables

The system presents in the following, non-dimensionalized form. This is not the original Bonheffer-Van-der-Pol model, however it is topologically equivalent and much easier to analyse.

$$\frac{dV}{dt} = V(\alpha + V)(1 - V) - W + z \tag{8}$$

$$\frac{dW}{dt} = \beta V - cW \tag{9}$$

In this model, V - in general - represents activation, and in an excitable tissue context represents voltage. and W - in general represents recovery, and has no realistic interpretation in a biological context, it would be the sum of all positive W.R.T.cell currents. There are three parameters in the model, α , β , c, and z. Of the three parameters, we have significant interest in z, as it will determine the bifurcation of the model. Neurologically, z represents the DC offset from experimentally injected current of the neuron under study. In our extension to this model, we will replace z with a gap junctional current from coupled cells (more in section 2). The additional parameters α , β , and c represent 'rate - like' constants which determine oscillator traits such as amplitude, frequency, duty cycle, and excitability.

1.2 Purpose of Model and Examination

This model, a simplification of the original Hodgkin Huxley model, intends to model spiking neurons. Tuning the individual parameters can adapt the model to fit various states and conditions which may be investigated by researchers. Various combinations of of parameters can also model various neuronal behaviors such as bursting and chattering. These models can be put together to form multi dimensional models of neuron trends. The state of the neuron relies on the z parameter. In this project, we plan to evaluate the stability of system at rest and with regards to the bifurcation parameter z. We plan to evaluate how the system responds and bifurcates with regards to this parameter, and interpret these stabilities biologically.

Additionally, we will examine and interpret the

- 1. Phase Plane
- 2. Bifurcation Diagram
- 3. Time Series

The model itself is already non-dimensionalized, though in our analysis we will provide a full description of the author's rationale. Additionally, we will fully classify the model according to the Haefner et. al criteria and justify our classification.

2 Modelling Myocardial Necrosis

2.1 Necrosis Propagation

As part of the normal cell cycle, programmed cell death (apoptosis) exists and it serves several purposes. In most cases, it is both irreversible and adaptive (beneficial) for the organism. However in some cases, apoptosis can occur accidentally in which case an irreversible destruction of the cell will occur. In our model, we assume that necrosis is a function of an underlying accumulation of pro-apoptotic proteins, potentially introduced as a byproduct of disease. As such, necrosis is a continuous state of each cell. For cell i of n, define pro-necrotic burden as ν_i , with total burden of the system equal to $\frac{\sum_{i=1}^{n}\nu_i}{n} = \frac{1}{n}\sum_{i=1}^{n}\nu_i$. Note that $\nu_1, ..., \nu_n \in [0,1]$ and is a monotonically increasing function with respect to time.

2.2 Investigation Goals and Expected Results

In order to model the heart and disease models, we will couple cardiac myocytes (with gap junctions) to form a one dimensional caricature of cardiac electrical propagation. Gap junctions act by physically linking 2 cells, i.e. the current will be proportional to the electrical potential between the 2 cells. As such, a spike in one cell can cause a spike in another cell. By having the final cell feedback onto the first cell, we can form simplistic model of the heart.

We want to ask how the temporal effects and distribution of cardiac death affects the overall frequency of the network. We expect that each diseased cell will act as a delay line, in which if 2 cells have equal states of disease α and β , this will be equal to a single cell with a disease state $\alpha + \beta$.

Define D, the delay in the enntire system caused by pro-necrotic built up, such that $D = T_n - T_t$ where T_r represents the period of the network without disease and T_t represents the period of the network at time t.

Consider a system with two, gap junctionally connected, neurons N_1 and N_2 . D is a function such that $D(N_1(\nu_1), N_2(\nu_2))$ defines the system's delay caused by ν_1 at N_1 and ν_2 at N_2 . It is our hypothesis:

$$\nu_1 = \alpha \tag{10}$$

$$\nu_2 = \beta \tag{11}$$

$$\nu_3 = \beta + \alpha \tag{12}$$

$$\nu_4 = 1 \tag{13}$$

$$D(N_1(\nu_1), N_2(\nu_2)) = D(N_1(\nu_3), N_2(\nu_4))$$
(14)

To examine and support our hypothesis, we will computationally model the above model using the FN model and gap junctionally coupled cardiac myocytes. We will construct graphics that demonstrate and illustrate our hypothesis.

Additionally, we will explore how the system responds in a variety of circumstances:

- 1. Does distance (Distance between N_i and $N_j = j i$) influence D
- 2. Is D decreased more by a few heavily burdened cells, or by a large number of moderately burdened cells
- 3. At what point does the build up of pro-necrotic proteins become a health issue (health being defined such that $T_t \geq kT_n; k \in [0,1]$)

The above will be examined in both a static (ν_1 remains contant with time) and a dynamic (ν_i is a function of t) model of cardiac necrosis.

2.3 Translational Perspective

The heart serves a vital role in many organisms. The timely and regular delivery of oxygenated blood is essential to survival, and thus any decrease from optimum efficiency is of consequence. Investigations into potential mechanistic explainations and modeling of this degradation are highly interesting in the context of both accute and chronic cardiac disease.

In particular, myocardial necrosis can be caused by accute oxygen deprivation of myocardial tissues, commonly experienced during a heart attack. Other potential etiologies stem from depriving this tissue of the blood it requires; dangerous increase in physical stresses, coronary artery disease, and accute hemerage all have the potential to play into cardiac necrosis, perhaps in tandem.

In our investigation into the decreasing efficiency of a heart-like system, we will model several scenarios that may be of consequence to sufferers of

cardiac necrosis. In particular, we study two cases. The first, static necrosis, closely resembles a short term or acute stress on myocardial tissue, be it a myocardial infarction or trauma. We will investigate how the distribution of necrosis affects the system as a whole through the FN model. Through a translation lens, this equates to the decrease in efficiency as a consequence of various kinds of trauma. The second case will be a slowly increasing level of necrosis. This more closely resembles coronary artery disease, which has the potential to starve the tissues of oxygen for many years, slowly leading to health deficits in older age. We study the approximate timeframe for simulated tachycardia, or dangerously low heart beat.

Throughout the investigation, we refer to a cell's necrosis burden as analagous to a protein build up, though this simply refers to the degree of degredation for that specific cell.

Our proposed extension to the FN model plans to shed light on the geneisis, progression, and finally the consequences of myocardial necrosis.

3 Discussion of Report

3.1 Proposed Format

We propose the following format for the final report:

- 1. Introduction
- 2. Primary Model Analysis
 - (a) Variable Description
 - (b) Model Classification
 - (c) Time series
 - (d) Phase Plane
 - (e) Orbit / Bifurcation Diagrams
- 3. Cardiac Necrosis Extension
 - (a) Description of extension
 - (b) Variable Explanation
 - (c) Time series with incorporation of ν
 - (d) Computational Model Rational
 - (e) Static ν

- i. Distance
- ii. Differential burden of System
- iii. Interpretation
- (f) Dynamic ν
 - i. Distance
 - ii. Differential burden of System
 - iii. Interpretation
- 4. Discussion
- 5. Conclusions and Further Work
- 6. References
- 7. Appendix I: Code

3.2 Timeline

We propose a checkup in approximately one week to assess our progress and make sure that the project presented is reasonable in scope for the course. If possible, we would like to present a draft to ensure that we are on the right path.

4 References

- 1. Abbott, L. . Lapicques introduction of the integrate-and-fire model neuron 1907. Brain Res. Bull. 50, 303304 1999.
- 2. Hodgkin, A. L., Huxley, A. F. A quantitative description of membrane current and its application to conduction and excitation in nerve. J. Physiol. 117, 50044 1952.
- 3. Haefner, J. W. Models of Systems. Modeling Biological Systems: Principles and Applications $Springer, 2005.\ doi:10.1007/0-387-25012-3_1$