Geometric understanding of local fluctuation distribution of conduction time in lined-up cardiomyocyte network in agarose-microfabrication multi-electrode measurement assay

Kazufumi Sakamoto 1,* , Shota Aoki 1, Yuhei Tanaka 1, Kenji Shimoda 1, Yoshitsune Hondo 1 and Kenji Yasuda 1

1 Department of Pure and Applied Physics, Graduate School of Advanced Science and Engineering, Waseda University, 3-4-1 Okubo, Shinjuku, Tokyo, 169-8555, Japan.

* Corresponding author: sk.060603@akane.waseda.jp
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[Diagram showing MEA Chip, MEA System, Measurement Record and Analysis PC, and an image of Cardiomyocyte Networks with a histogram of conduction time between specific two electrodes.]
Abstract

We examined characteristics of the propagation of conduction in width-controlled cardiomyocyte cell networks for understanding the contribution of the geometrical arrangement of cardiomyocytes for their local fluctuation distribution. We tracked a series of extracellular field potentials of linearly lined-up mouse primary cardiomyocytes and human ES cardiomyocytes with 100 kHz sampling intervals of multi-electrodes signal acquisitions and an agarose microfabrication technology. Conduction time between two neighbor microelectrodes showed Gaussian distribution, which indicates this conduction propagation in a unit length was a stochastic firing phenomenon. However, the distributions of conduction time were not expanded but maintained within an identical range of distribution regardless of their propagation distances from 0.3 mm to 1.5 mm, which is against the expected distance-dependent enlarging of the distribution based on the faster firing regulation. In contrast, when Quinidine was applied to the cardiomyocytes, the distributions of conduction time were expanded as propagation distance increased as predicted by the conduction propagation model of faster firing regulation. The results indicate the “faster firing regulation” is not sufficient to explain this conservation of the propagation time distribution in cardiomyocyte networks.

Keywords

on-chip cell network assay, multi microelectrode array, external field potential measurement, conduction distribution, cardiomyocyte network
Can explain the excitation conduction by the faster firing regulation?

Simulation of conduction time distribution in the faster firing regulation model

The distribution of conduction time dispersed as the firing signal propagated.

Fig.1. The histogram of conduction time distribution.
Result and Discussion (1)

Agarose Microfabrication and Cell Culture

Fig.2. MEA chip microfabrication procedure.

Fig.3. Agarose microfabrication technology.

Fig.4. Cultured cardiomyocytes on MEA chip.
Result and Discussion (2)

Measurement System and Data Analysis

Fig. 5. MEA system.

Fig. 6. Conduction time measurement and analysis.

MEA chip

Amplifier

AD converter

Computer

incubated at 37°C, 5% CO₂

MEA chip

Amplification × 5000

Analog-to-digital conversion 100 kHz

Recording

Field Potential

Inter-spike interval

Cardiomyocyte network

Conduction Time k

Conduction Time k+1

300 μm

Potential

Time
Result and Discussion (3)

Distribution of Conduction Time

- Mouse Primary Cardiomyocyte Networks

Fig. 7. Primary cardiomyocyte networks.

Fig. 8. Histograms of conduction time between specific two electrodes.
Human ES Cardiomyocyte Networks

Fig. 9. Histograms of conduction time.

Effect of conduction control compound

Fig. 10. Histograms of conduction time when 3 µM Quinidine was applied.
Result and Discussion

Human ES Cardiomyocyte Networks

The distributions of conduction time maintained within an identical range of distribution regardless of their propagation distances.

The distributions of conduction time were expanded as propagation distance increased.
Result and Discussion (5)

Distribution of Conduction Time

Fig11. Comparison of fluctuations of conduction time in cardiomyocyte networks.
Can explain the excitation conduction by the faster firing regulation?

Simulation Result

Experimental Result of 300 µm width hES cardiomyocyte network
Result and Discussion (6)

Can explain the excitation conduction by the faster firing regulation?

Simulation Result

Experimental Result of 300 µm width hES cardiomyocyte network
Conclusions

(1) We succeed in constructing measurement assay of excitation conduction time in width-controlled linearly lined-up cardiomyocyte network on the multi electrode array chip.

(2) We observed the distributions of conduction time maintained their range of distribution without any expansion regardless of its propagation distances from 0.3 mm up to 1.5 mm, which is against the conventional conduction connection rule, “faster firing regulation”.

(3) We also observed the distributions of conduction time were expanded as propagation distance increased when Quinidine was applied, which was followed to “the faster firing regulation.”

Above (2) and (3) suggest the existence of some unknown cooperative conduction propagation regulation in cardiomyocyte conduction, which is disappeared by the sodium channel blocking.

In detail, please visit our publication in this issue Sakamoto, Kazufumi, et al. *Micromachines* 11.12 (2020): 1105.
Acknowledgments

This research was funded by research and development projects of the Industrial Science and Technology Program, the New Energy and Industrial Technology Development Organization (NEDO, P08030), JSPS KAKENHI Grant Number JP17H02757, JST CREST program, and Waseda University Grant for Special Research Projects (2016S-093, 2017B-205, 2017K-239, 2018K-265, 2018B-186, 2019C-559), the Leading Graduate Program in Science and Engineering for Waseda University from MEXT, Japan.

We thank to Dr. Akihiro Hattori, Mr. Masao Odaka and all the members of Yasuda lab for their technical supports, discussion, and suggestions.