Key words: mitochondrial network dynamics, ROS, RIRR, superoxide anion (O$_2^-$), hydrogen peroxide (H$_2$O$_2$), oxidative stress, ABM

Abbreviations: ABM, agent-based modeling; RIRR, ROS-induced ROS release; ROS, reactive oxygen species; SOD, superoxide dismutase

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*Correspondence to: Chulhee Choi; Email: cchoi@kaist.ac.kr

Oxidative stresses can induce rapid depolarization of inner mitochondrial membrane potential and subsequent impairment of oxidative phosphorylation. Damaged mitochondria produce more reactive oxygen species (ROS), particularly the superoxide anion (O$_2^-$) and hydrogen peroxide (H$_2$O$_2$), which potentiate mitochondria-driven ROS propagation, so-called ROS-induced ROS release (RIRR), via activation of an inter-mitochondrial signaling network. In this context, mitochondrial network dynamics, such as their density, number, and spatial distribution, can affect mitochondria-driven ROS propagation. To investigate this inter-mitochondrial communication, we developed a mathematical model using an agent-based modeling approach and tested the effect of mitochondrial network dynamics on RIRR for mitochondria under various conditions. Simulation results show that mitochondrial network dynamics are critical determinants of inter-mitochondrial ROS signaling patterns and main messenger ROS molecules. We further elucidated the potential mechanism of these actions, which is conversion of major messenger molecules involved in ROS signaling. Collectively, we propose that mitochondrial network dynamics can determine cellular responses to oxidative stress by switching the molecular species involved in cellular signaling.

Among the many intracellular organelles, mitochondria are the powerhouse that provides energy for other cellular processes. During cellular respiration, reactive oxygen species (ROS) are inevitably produced, which can cause oxidative damage to many cellular components including proteins, nucleic acids, lipid membranes, and even the mitochondria themselves. Damaged mitochondria produce more ROS, particularly the superoxide anion (O$_2^-$) and hydrogen peroxide (H$_2$O$_2$), which potentiate mitochondria-driven ROS propagation, called ROS-induced ROS release (RIRR), via activation of an inter-mitochondrial signaling network (Fig. 1A). Several studies have reported on RIRR and how loss of function of a small number of mitochondria might influence overall cell functioning. However, most previous RIRR studies were based on a specialized context of intracellular mitochondrial dynamics within cardiomyocytes, taking advantage of the electrically independent distribution of rigid, lattice-like mitochondria suitable for high precision confocal line-scan imaging. In the real world, mitochondrial network dynamics, such as their density, number, and spatial distribution, can affect the pattern of ROS propagation in cells. Thus, many unknowns regarding mitochondrial dynamics and mitochondria-driven ROS propagation remain, and some current experimental results do not fit the existing model. To resolve this problem, a more systemic approach to study inter-mitochondrial networks is required.
We created a mathematical model of inter-mitochondrial communication using the agent-based modeling (ABM) method, which allows controlling many variables of spatial information in molecular signaling systems. A mathematical and computational approach based on ABM is meaningful and reasonable in that it can simulate diverse scenarios by controlling variables independently, whereas design of in vitro/in vivo experiments incorporating such variables is practically impossible. In our model, local introduction of initial oxidative stress causes damage in that area, and ROS are propagated in every cycle. Simulation results revealed that the response to oxidative stress differed according to the status of mitochondrial network dynamics. The cardiomyocyte model was more dependent on the initial dose of oxidative stress compared with irregular distribution or low mitochondrial density models. Hence, an identical oxidative stress input can cause different responses, and the eventual fate of cells is related to mitochondrial network dynamics. We also elucidated conversion of main messenger ROS molecules from $O_2^\cdot$ to $H_2O_2$ in the irregularly distributed and low-number mitochondria models (Fig. 1B–D). Our findings clearly suggest that mitochondrial network dynamics may determine the intracellular ROS propagation profile and cellular responses to oxidative stress by switching inter-mitochondrial ROS messengers.

In previous research, $O_2^\cdot$, which is a more reactive ROS species than $H_2O_2$, has been considered important in the process of RIRR. Because such research was performed using only the cardiomyocyte model, which has a strictly regular distribution and high number of mitochondria, this explanation is reasonable only in that specific system. However, our simulation results revealed that mitochondrial network dynamics determine the identity of inter-mitochondrial ROS signaling and that $H_2O_2$-dependent ROS propagation occurs according to the distribution of mitochondria and distance between neighboring mitochondria.

With this knowledge, several extraordinary findings in various cell types can be explained in that inhibition or reduction of CuZn-superoxide dismutase (SOD) increases resistance to oxidative stress, and that treatment of cells with the SOD mimic MnTBAP triggers abnormal hyperpolarization of mitochondria under oxidative stress (unpublished data). Our data also provide a plausible explanation for the correlation between oxidative stress vulnerability and the status of the cytoskeleton in cardiomyocytes. Colchicine is reported to disrupt microtubules by binding to tubulin subunits. Because mitochondria are attached to and move along these microtubules, treatment with colchicine may break up the regular distribution of mitochondria into an irregular network. Therefore, our results also provide an explanation of how colchicine attenuates ischemic preconditioning-induced cardioprotection.

We showed that the degree of ROS propagation, effect of each antioxidant enzyme and vulnerability to oxidative stress deciding the ultimate fate of a cell can be changed depending on the characteristics of mitochondrial network dynamics. What would be the cellular advantage of determining the profile of ROS signaling by mitochondrial network dynamics? Mitochondria have previously been shown to be highly dynamic organelles whose morphology, distribution, and activity can be regulated by fusion, fission, and migration. Thus, cells can modulate their own mitochondrial network dynamics to efficiently adapt to environmental oxidative stresses. By introducing ABM-based computational analysis, the complex dynamics of various cellular behaviors regarding RIRR was better understood and it may provide a rational background for development of mechanism-based therapeutic interventions.

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