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Integrative Modeling of Mitochondrial Metabolism

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1 Introduction

We are in the process of constructing a kinetic model of mitochondrial metabolism including the electron transport (respiratory chain), the TCA cycle, the fatty acid metabolism (β oxidation), the inner-membrane metabolite carriers, the protein carriers, and the mitochondrial gene expression system. In this work, we report a model of these metabolic pathways that has been recently completed using the E-CELL system [2], a generic simulation environment for cellular simulation based on multiple ordinary differential equations.

2 Methods and Results

The model consists of the following number of enzymatic reactions: six (6) for the respiratory chain, eight (8) along with some ancillary reactions for the TCA cycle, seven (7) for the β oxidation, seven (7) for the metabolite carrier system, one (1) for the protein carrier system, and five (5) enzymatic reactions and several accompanying reactions for the mitochondrial gene expression system (Table 1). All of the enzymatic reactions are modeled based on kinetic parameters found in the literature or estimated by various parameter-tuning methods [1].

Figure 1 presents an example run of E-CELL from an *in silico* experiment conducted using this model. The figure shows the temporal transition of the concentration of ubiquinone and ubiquinol, a pair of electron transporters; the former is transformed into the latter by accepting two electrons. At the point of 70 seconds, the TCA cycle and the β oxidation pathway were disrupted in order for us to observe their influence on the respiratory chain. Consequently, a sudden change of the ubiquinone/ubiquinol ratio took place (Figure 1).

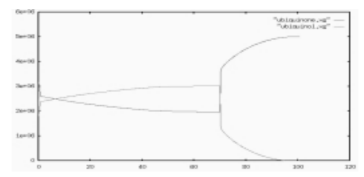


Figure 1: Temporal transition of ubiquinone and ubiquinol

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This transition occurs because ETF-Q oxidoreductase in the β oxidation pathway and Succinate dehydrogenase in the TCA cycle had the ability to reduce ubiquinone into ubiquinol. Hence, the example run shows that the TCA cycle and the β oxidation including those enzymes play an important role in maintaining the rate of electron transport.

We will continue to work on this mitochondrial model, with the eventual goal of applying it to pathological analyses of mitochondrial diseases such as Leigh syndrome and mitochondrial myopathy, encephalopathy, lactic acidosis, and stroke-like episodes (MELAS).

Group	Enzyme	Group	Enzyme
respiratory chain	NADH dehydrogenase Ubiquinol:Cytochrome c oxidoreductase Cytochrome c oxidase ATP synthase	β oxidation	Acyl-CoA dehydrogenase Enoyl-CoA hydratase 3-hydroxyacyl-CoA dehydrogenase 3-ketoacyl-CoA thiolase
TCA cycle	Pyruvate dehydrogenase complex Malic enzyme Pyryvate carboxylase		ETF:Q oxidoreductase CPT I CPT II
	Citrate synthase Citrate hydratase Isocitrate dehydrogenase 2-oxoglutare dehydrogenase complex Succinyl-CoA synthetase Succinate dehydrogenase Fumarase	Metabolite carriers	Pyruvate carrier Citrate carrier Dicarboxylate carrier Oxoglutarate carrier Adenine nucleotide carrier Phosphate carrier Carnitine carrier
	Malate dehydrogenase	Protein carrier mitochondrial gene expression system	Protein carrier RNA polymerase mtRNase P Aminoacyl-tRNA synthetase Peptidyl transferase MPP

Table 1: The reactions in the model

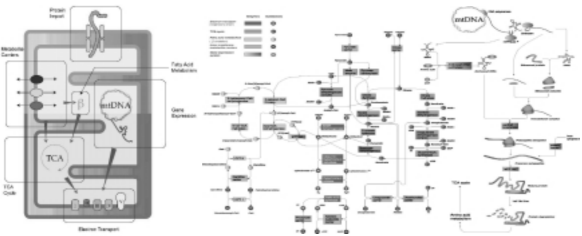


Figure 2: a) An overview of the model and b) the modeled reactions

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