

SELF-ORGANIZATION AND FRACTALITY IN A METABOLIC PROCESSES OF THE KREBS CYCLE

V. I. GRYTSAY^{1,*}, I. V. MUSATENKO²

¹*Bogolyubov Institute for Theoretical Physics,
National Academy of Sciences of Ukraine, Kyiv;
e-mail: vgrytsay@bitp.kiev.ua;*

²*Taras Shevchenko National University of Kyiv, Ukraine;
e-mail: ivmusatenko@gmail.com*

The metabolic processes of the Krebs cycle is studied with the help of a mathematical model. The autocatalytic processes resulting in both the formation of the self-organization in the Krebs cycle and the appearance of a cyclicity of its dynamics are determined. Some structural-functional connections creating the synchronism of an autoperiodic functioning at the transport in the respiratory chain and the oxidative phosphorylation are investigated. The conditions for breaking the synchronization of processes, increasing the multiplicity of cyclicity, and for the appearance of chaotic modes are analyzed. The phase-parametric diagram of a cascade of bifurcations showing the transition to a chaotic mode by the Feigenbaum scenario is obtained. The fractal nature of the revealed cascade of bifurcations is demonstrated. The strange attractors formed as a result of the folding are obtained. The results obtained give the idea of structural-functional connections, due to which the self-organization appears in the metabolism running in a cell. The constructed mathematical model can be applied to the study of the toxic and allergic effects of drugs and various substances on cell metabolism.

Key words: Krebs cycle, metabolic processes, self-organization, fractality, strange attractor, Feigenbaum scenario.

The search for general physical laws of self-organization in the Nature is one of the most important problems of natural sciences. As the principal problem, we mention the study of the nature of life, namely how the catalyzed enzymatic reactions create the internal space-time ordering of a cell. The development of mathematical models of such systems allows one to obtain a lot of important information.

Among various metabolic processes, the Krebs cycle is common for all cells [1]. The cycle involving tricarboxylic acids occupies the principal place, to which practically all metabolic paths lead. It is the final point of the catabolism of a "cell fuel," the central place of a cell respiration, and the source of molecules-predecessors, from which the aminoacids, carbohydrates, fat acids, and other compounds important for the functioning of cells are then synthesized. Its study will allow one to find the general regularities of the operation of a cell.

The functioning and the regulation of the Krebs cycle was studied experimentally and theoretically [2-10] within a restricted number of mathematical models. Many of the models considered only the stoichiometry and segments of the cycle. As the main difficulty, the deficit of experimental data should be mentioned.

It is difficult to study separately this cycle without the consideration of the functioning of a cell as a whole, since it is impossible to determine the internal parameters of a metabolic processes without the knowledge of the external parameters of the medium, where a cell lives. Our studies are based on the mathematical model of the unstable growth of *Candida utilis* cells on ethanol, which was developed by Professor V. P. Gachok [11, 12] in view of the experimental data published in [13]. Analogous problems, namely, the modeling of a cell growth, were considered by J. Monod, V. S. Podgorskyi, L. N. Drozdov-Tikhomirov, N. T. Rakhimova, G. Yu. Riznichenko and others [14-18].

Our refined mathematical model involves the formation of carbon dioxide and considers its influence on the kinetics of a metabolic process.

In the present work, we study the unstable modes of the cultivation of cells and will find the parameters and structural-functional connections of the metabolic process running in a cell that admit the appearance of oscillations. We consider the intracellular autooscillations characteristic of the metabolic process on the level of redox reactions of the Krebs cycle. These autooscillations are related to the process cyclicity and characterize the self-organization inside a cell.

Analogous oscillatory modes are observed in the processes of photosynthesis and glycolysis, a variation in the concentration of calcium in a cell, oscillations in the heart muscle, and some biochemical processes [19–27].

We note that the description of the metabolic process within the presented model is not based on the use of conditional reagents, as was made earlier in other models, but involves the real intracellular components, which will allow us to find an actual mechanism of self-organization.

Mathematical Model

The general scheme of the process is presented in Fig. 1. According to it with regard for the mass balance, we have constructed the mathematical model given by Eqs. (1)–(19).

$$\frac{dS}{dt} = S_0 \frac{K}{K + S + \gamma\psi} - k_1 V(E_1) \frac{N}{K_1 + N} V(S) - \alpha_1 S, \quad (1)$$

$$\frac{dS_1}{dt} = k_1 V(E_1) \frac{N}{K_1 + N} V(S) - k_2 V(E_2) \frac{N}{K_1 + N} V(S_1), \quad (2)$$

$$\frac{dS_2}{dt} = k_2 V(E_2) \frac{N}{K_1 + N} V(S_1) - k_3 V(S_2^2) V(S_3) - k_4 V(S_2) V(S_8), \quad (3)$$

$$\frac{dS_3}{dt} = k_4 V(S_2) V(S_8) - k_3 V(N^2) V(S_3^2) - k_3 V(S_2^2) V(S_3), \quad (4)$$

$$\frac{dS_4}{dt} = k_3 V(N^2) V(S_3^2) - k_7 V(N) V(S_4) - k_8 V(N) V(S_4), \quad (5)$$

$$\frac{dS_5}{dt} = k_7 V(N) V(S_4) - 2k_9 V(L_1 - T) V(S_5), \quad (6)$$

$$\frac{dS_6}{dt} = 2k_9 V(L_1 - T) V(S_5) - k_{10} V(N) \frac{S_6^2}{S_6^2 + 1 + M_1 S_8}, \quad (7)$$

$$\frac{dS_7}{dt} = k_{10} V(N) \frac{S_6^2}{S_6^2 + 1 + M_1 S_8} - k_{11} V(N) V(S_7) - k_{12} \frac{S_7^2}{S_7^2 + 1 + M_2 S_9} V(\psi^2) + k_3 V(S_2^2) V(S_3), \quad (8)$$

$$\frac{dS_8}{dt} = k_{11} V(N) V(S_7) - k_4 V(S_2) V(S_8) + k_6 V(T^2) \frac{S^2}{S^2 + \beta_1} \cdot \frac{N_1}{N_1 + (S_5 + S_7)^2}, \quad (9)$$

$$\frac{dS_9}{dt} = k_{12} \frac{S_7^2}{S_7^2 + 1 + M_2 S_9} V(\psi^2) - k_{14} \frac{XTS_9}{(\mu_1 + T)[(\mu_2 + S_9 + X + M_3(1 + \mu_3\psi))]S}, \quad (10)$$

$$\frac{dX}{dt} = k_{14} \frac{XTS_9}{(\mu_1 + T)[(\mu_2 + S_9 + X + M_3(1 + \mu_3\psi))]S} - \alpha_2 X, \quad (11)$$

$$\frac{dQ}{dt} = -k_{15} V(Q) V(L_2 - N) + 4k_{16} V(L_3 - Q) V(O_2) \frac{1}{1 + \gamma_1 \psi^2}, \quad (12)$$

$$\frac{dO_2}{dt} = O_{20} \frac{K_2}{K_2 + O_2} - k_{16} (L_3 - Q) V(O_2) \frac{1}{1 + \gamma_1 \psi} - k_8 V(N) V(S_4) - \alpha_3 O_2, \quad (13)$$

$$\frac{dN}{dt} = -k_7 V(N) V(S_4) - k_{10} V(N) \frac{S_6^2}{S_6^2 + 1 + M_1 S_8} - k_{11} V(N) V(S_7) - k_3 V(N^2) V(S_3^2) + k_{15} V(Q) V(L_2 - N) - k_2 V(E_2) \frac{N}{K_1 + N} V(S_1) - k_1 V(E_1) \frac{N}{K_1 + N} V(S), \quad (14)$$

$$\frac{dT}{dt} = k_{17} V(L_1 - T) V(\psi^2) + k_9 V(L - T) V(S_3) - \alpha_4 T - k_{18} k_6 V(T^2) \frac{S^2}{S^2 + \beta_1} \cdot \frac{N_1}{N_1 + (S_5 + S_7)^2} - k_{19} k_{14} \frac{XTS_9}{(\mu_1 + T)[\mu_2 + S_9 + X + M_3(1 + \mu_3\psi)]S}, \quad (15)$$

$$\frac{d\psi}{dt} = 4k_{15} V(Q) V(L_2 - N) + 4k_{17} V((L_1 - T) V(\psi^2) - 2k_{12} \frac{S_7^2}{S_7^2 + 1 + M_2 S_9} V(\psi^2) - \alpha\psi, \quad (16)$$

$$\frac{dE_1}{dt} = E_{10} \frac{S^2}{\beta_2 + S^2} \frac{N_2}{N_2 + S_1} - n_1 V(E_1) \frac{N}{K_1 + N} V(S) - \alpha_5 E_1, \quad (17)$$

$$\frac{dE_2}{dt} = E_{20} \frac{S_1^2}{\beta_3 + S_1^2} \frac{N_3}{N_3 + S_2} - n_2 V(E_2) \frac{N}{K_1 + N} V(S_1) - \alpha_6 E_2, \quad (18)$$

$$\frac{dC}{dt} = k_8 V(N) V(S_4) - \alpha_7 C, \quad (19)$$

where $V(X) = X/(1 + X)$ is the function that describes the adsorption of the enzyme in the region of local coupling. The variables of the system are dimensionless.

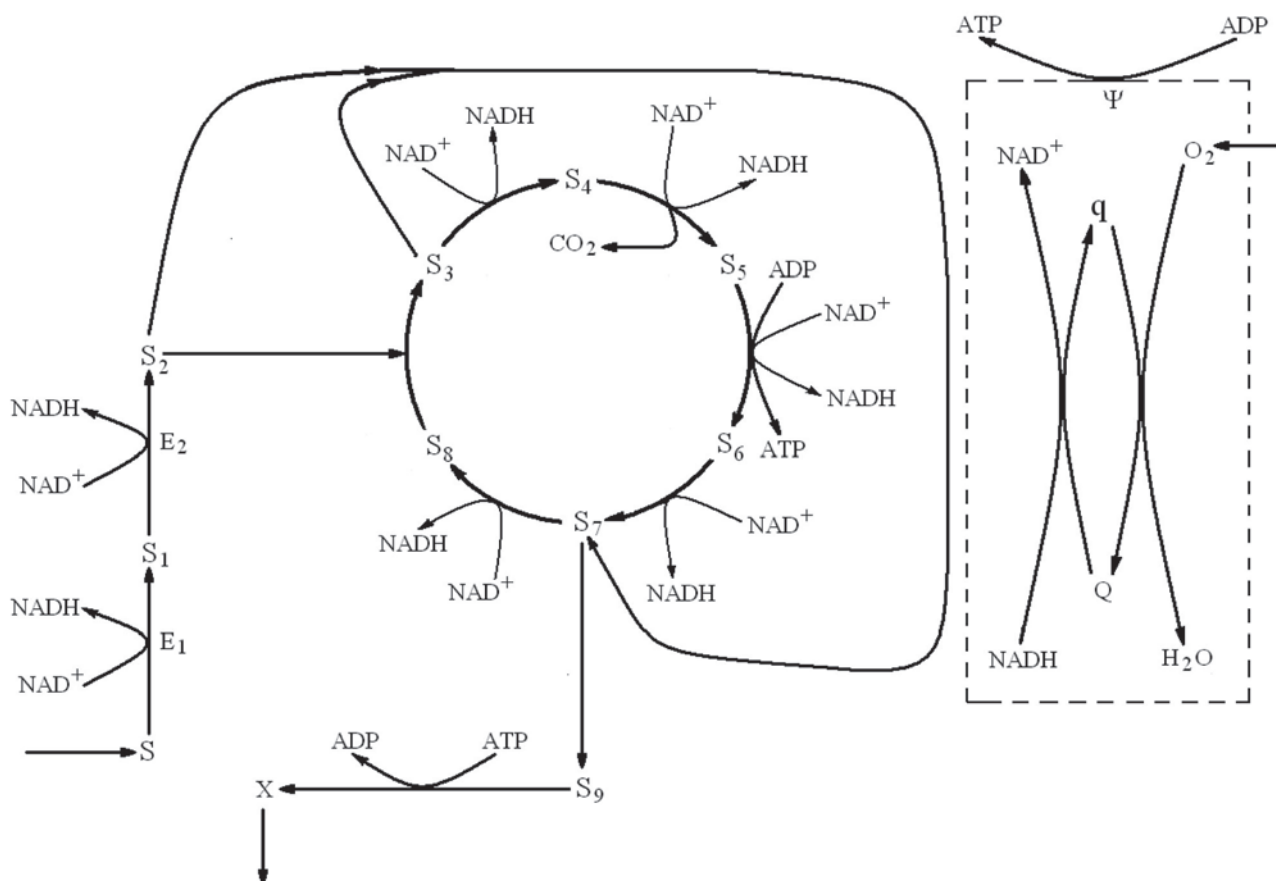


Fig. 1. General scheme of the metabolism of growth of cells *Candida utilis* on ethanol [12]

The internal parameters of the system are as follows: $k_1 = 0.3$; $k_2 = 0.3$; $k_3 = 0.2$; $k_4 = 0.6$; $k_5 = 0.16$; $k_6 = 0.7$; $k_7 = 0.08$; $k_8 = 0.022$; $k_9 = 0.1$; $k_{10} = 0.08$; $k_{11} = 0.08$; $k_{12} = 0.1$; $k_{14} = 0.7$; $k_{15} = 0.27$; $k_{16} = 0.18$; $k_{17} = 0.14$; $k_{18} = 1$; $k_{19} = 10$; $n_1 = 0.07$; $n_2 = 0.07$; $L = 2$; $L_1 = 2$; $L_2 = 2.5$; $L_3 = 2$; $K = 2.5$; $K_1 = 0.35$; $K_2 = 2$; $M_1 = 1$; $M_2 = 0.35$; $M_3 = 1$; $N_1 = 0.6$; $N_2 = 0.03$; $N_3 = 0.01$; $\mu_1 = 1.37$; $\mu_2 = 0.3$; $\mu_3 = 0.01$; $\gamma = 0.7$; $\gamma_1 = 0.7$; $\beta_1 = 0.5$; $\beta_2 = 0.4$; $\beta_3 = 0.4$; $E_{1_0} = 2$; $E_{2_0} = 2$.

The external parameters determining the flow-type conditions are chosen as $S_0 = 0.05055$; $O_{2_0} = 0.06$; $\alpha = 0.002$; $\alpha_1 = 0.02$; $\alpha_2 = 0.004$; $\alpha_3 = 0.01$; $\alpha_4 = 0.01$; $\alpha_5 = 0.01$; $\alpha_6 = 0.01$; $\alpha_7 = 0.0001$.

The model covers the processes of substrate-enzymatic oxidation of ethanol to acetate, cycle involving tri- and dicarboxylic acids, glyoxylate cycle, and respiratory chain. Model (1)–(19) is improved as compared with the model used in [11, 12], since it involves the formation of CO_2 in the Krebs cycle, which affects the running of the metabolic process. Some parameters of our model are taken from [11, 12].

The incoming ethanol S is oxidized by the alcohol dehydrogenase enzyme E_1 to acetaldehyde S_1 (1) and then by the acetal dehydrogenase enzyme E_2 to acetate S_2 (2), (3). The formed acetate can participate in the cell metabolism and can be exchanged with the environment. The model accounts for this situation by the change of acetate by acetyl-CoA. On the first stage of the Krebs cycle due to the citrate synthase reaction, acetyl-CoA jointly with oxalacetate S_8 formed in the Krebs cycle produce citrate S_3 (4). Then substances S_4 – S_8 are created successively on stages (5)–(9). In the model the Krebs cycle is represented by only those substrates which participate in the reduction of NADH and the phosphorylation $\text{ADP} \rightarrow \text{ATP}$. Acetyl-CoA passes along the chain to malate represented in the model as intramitochondrial S_7 (8) and cytosolic S_9 (10) ones. Malate can be also synthesized in another way related to the activity of two enzymes: isocitrate lyase and malate synthetase. The former catalyzes the splitting of isocitrate to succinate, and the latter catalyzes the condensation of acetyl-CoA with glyoxylate and formation of malate. This glyoxylate-linked way is

shown in Fig. 1 as an enzymatic reaction with the consumption of S_2 and S_3 and formation of S_7 . The parameter k_3 controls the activity of the glyoxylate-linked way (3), (4), (8). The yield of S_7 into cytosol is controlled by its concentration, which can increase due to S_9 , by causing the inhibition of its transport with participation of protons of the mitochondrial membrane.

The formed malate S_9 is used by a cell for its growth, namely for the biosynthesis of protein X (11). The energy consumption of the given process is supported by the process $\text{ATP} \rightarrow \text{ADP}$. The presence of ethanol in the external solution causes the "ageing" of external membranes of cells, which leads to the inhibition of this process. The inhibition of the process also happens due to the enhanced level of the kinetic membrane potential ψ . The parameter μ_0 is related to the lysis and wash-out of cells.

In the model, the respiratory chain of a cell is represented in two forms: oxidized, Q , (12) and reduced, q , ones. They obey the integral of motion $Q(t) + q(t) = L_3$.

The oxygen concentration change in the respiratory chain is determined by Eq. (13).

The activity of the respiratory chain is affected by the level of NADH (14). Its high concentration leads to the enhanced endogenic respiration in the reducing process in the respiratory chain (parameter k_{15}). The accumulation of NADH occurs as a result of the reduction of NAD^+ at ethanol transformation and in the Krebs cycle. These variables obey the integral of motion $\text{NAD}^+(t) + \text{NADH}(t) = L_2$.

In the respiratory chain and the Krebs cycle, the substrate-linked phosphorylation of ADP with the formation of ATP (15) is also realized. The energy consumption due to the process $\text{ATP} \rightarrow \text{ADP}$ induces the biosynthesis of components of the Krebs cycle (parameter k_{18}) and the growth of cells on the substrate (parameter k_{19}). For these variables, the integral of motion $\text{ATP}(t) + \text{ADP}(t) = L_1$ holds. Thus, the level of ATP produced in the redox processes in the respiratory chain $\text{ADP} \rightarrow \text{ATP}$ determines the intensity of the Krebs cycle and biosynthesis of protein.

In the respiratory chain, the kinetic membrane potential ψ (16) is created under the running of reducing processes $Q \rightarrow q$. It is consumed at the substrate-linked phosphorylation $\text{ADP} \rightarrow \text{ATP}$ in the respiratory chain and the Krebs cycle. Its enhanced level inhibits the biosynthesis of protein and process of reduction of the respiratory chain.

Equations (17) and (18) describe the activity of enzymes E_1 and E_2 , respectively. We consider their biosynthesis (E_{10} and E_{20}), the inactivation

in the course of the enzymatic reaction (n_1 and n_2), and all possible irreversible inactivations (α_5 and α_6).

Equation (19) is related to the formation of carbon dioxide. Its removal from the solution into the environment (α_c) is taken into account. Carbon dioxide is produced in the Krebs cycle (5). In addition, it squeezes out oxygen from the solution (13), by decreasing the activity of the respiratory chain.

The study of system (1)–(19) was carried out with the help of the Runge-Kutta-Merson method. The analysis of modes was performed when the system approaches the asymptotic trajectory of an attractor. To this end, the calculations started after 1,000,000 preliminary steps. The solution accuracy was set to be 10⁻¹².

In our studies, we used the classical tools of nonlinear dynamics [28, 29] and some methods of simulation of biochemical processes developed by the authors in [30–35].

Results of Studies

According to the constructed mathematical model (1)–(19), it is possible to separate two autocatalytic processes creating the space-time self-organization of the mutually connected metabolic processes of the Krebs cycle and the respiratory chain.

The continuous operation of the Krebs cycle requires the permanent supply of acetyl-CoA into the cycle. This substance is formed in the catabolism of ethanol, being the "cell fuel." Then acetyl-CoA is condensed with oxalocitrate and forms citric acid. After the dehydrogenation and two decarboxylations, citric acid is gradually transformed along a circle ($S_3 \rightarrow S_8$) to CO_2 and H_2O . In this case, the regeneration of oxalocitrate occurs at the end. On each stage of this cycle, there occurs the reduction of co-enzyme NAD^+ to NADH. The reverse process of oxidation $\text{NADH} \rightarrow \text{NAD}^+$ is realized in the system of carriers of electrons in the respiratory chain (1st autocatalytic cycle).

The energy released as a result of the oxidation of acetyl-CoA is concentrated to a significant degree in macroergic phosphate bonds of ATP. This happens at the transfer of electrons from the Krebs cycle to the respiratory chain, where the oxidative phosphorylation $\text{ADP} \rightarrow \text{ATP}$ occurs. The formed ATP is consumed for the growth of cells and in the Krebs cycle itself $\text{ATP} \rightarrow \text{ADP}$ (2nd autocatalytic cycle).

The preservation of the functioning of the Krebs cycle depends of the key parameters of the system relating to the inverse paths of the reduction of oxalocitrate. They are the parameters k_{15} and

k_6 . The former determines the rate of oxidation, $\text{NADH} \rightarrow \text{NAD}^+$, in the respiratory chain and depends on its parameters: $Q(t) + q(t) = L_3$. The rate of the whole autocatalytic process $\text{NADH} \leftrightarrow \text{NAD}^+$ is determined by the rate of the redox process in the respiratory chain itself $Q \leftrightarrow q$.

The parameter k_6 determines the influence of ADP concentration on the rate of the metabolic process of the Krebs cycle.

Thus, the metabolic process in the Krebs cycle is characterized by the appearance of autooscillations, whose frequency and stability depend on the synchronization of the 1st and 2nd autocatalytic cycles. Let us consider the kinetics of autooscillations in the metabolic process of the Krebs cycle, which arise in its cyclic mode. We will study the stability of the Krebs cycle on the whole.

During one turnover of a cycle, one molecule of acetyl-CoA is completely oxidized to malate, and a new molecule of acetyl-CoA is formed at the input. In such a way, the continuous process of operation of the Krebs cycle is running and manifests the autooscillatory character. In Fig. 2 we show the kinetics of autooscillations of the cycle components. At $k_{15} = 0.23$, a 1-fold periodic cycle appears in the system. The oscillations of components occur synchronously with phase shifts corresponding to the space-time positions of these components in the cycle. The oscillations at the same frequency will occur in all components of the given metabolic process.

In Fig. 3, *a*, we show a phase-parametric diagram presenting the dependence of the multiplicity

of autooscillations of the input substrate, ethanol (S), on the parameter k_{15} . To construct the phase-parametric diagram, we used the method of cutting. In the phase space of trajectories of the system, we placed a cutting plane at $S_2 = 0.8$. If the trajectory crosses this plane in a certain direction, we indicate the value of chosen variable (S , in this case) on the phase-parametric diagram. It is seen from the phase-parametric diagram that at $k_{15}^j = 0.2328$, the doubling of the period of oscillations arises. At $k_{15}^{j+1} = 0.2483$, we observe the repeated period doubling. Then, at $k_{15}^{j+2} = 0.2516$, the period of autooscillations is doubled once more.

Let us separate a small part of the diagram with $k_{15} \in (0.25, 0.254)$ (Fig. 3, *a*) and magnify it (Fig. 3, *b*). This part of the phase-parametric diagram is identical to the whole diagram in Fig. 3, *a*. This indicates that, after the $(j + 2)$ th bifurcation, the next bifurcation arises, and so on to $(j + n)$. In other words, we have a cascade of bifurcations. At the following decrease in the scale of the diagram, the pattern will repeat up to the critical value of the parameter k_{15} , after which we observe the appearance of a deterministic chaos. For the derived sequence of bifurcations, we have the relation

$$\lim_{i \rightarrow \infty} \frac{k_{15}^{j+1} - k_{15}^j}{k_{15}^{j+2} - k_{15}^{j+1}} \approx 4.696969\dots$$

This number is very close to the Feigenbaum universal constant $\delta = 4.669211660910\dots$, which characterizes the infinite cascade of bifurcations at the transition to a deterministic chaos. Thus, the part under consideration is characterized by

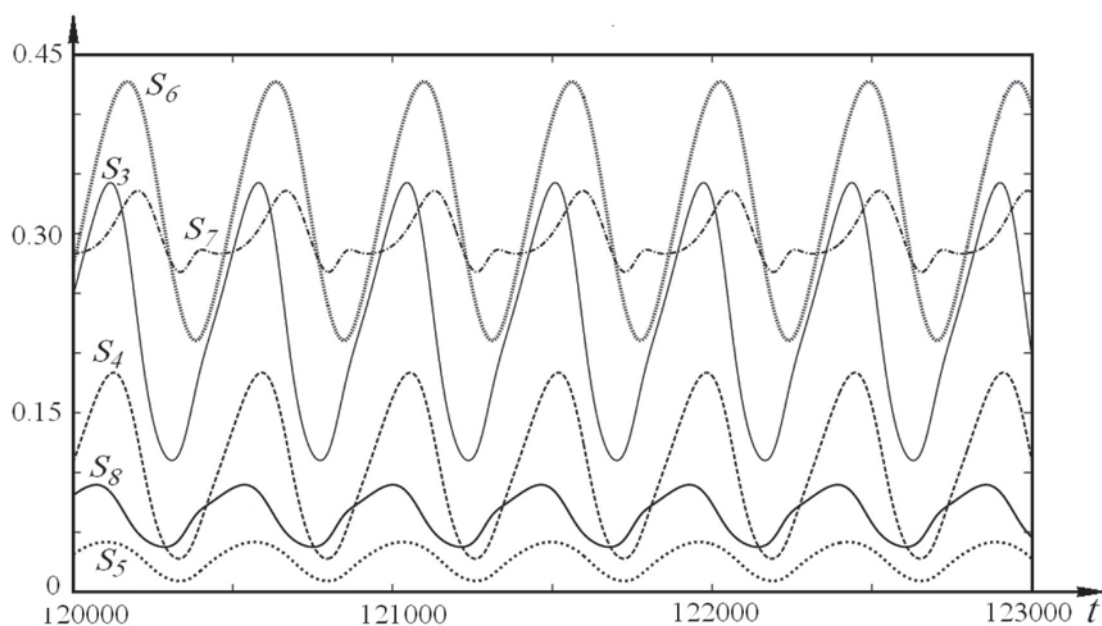


Fig. 2. Kinetics of autooscillations of components of the Krebs cycle $S_3 - S_8$, at $k_{15} = 0.23$.

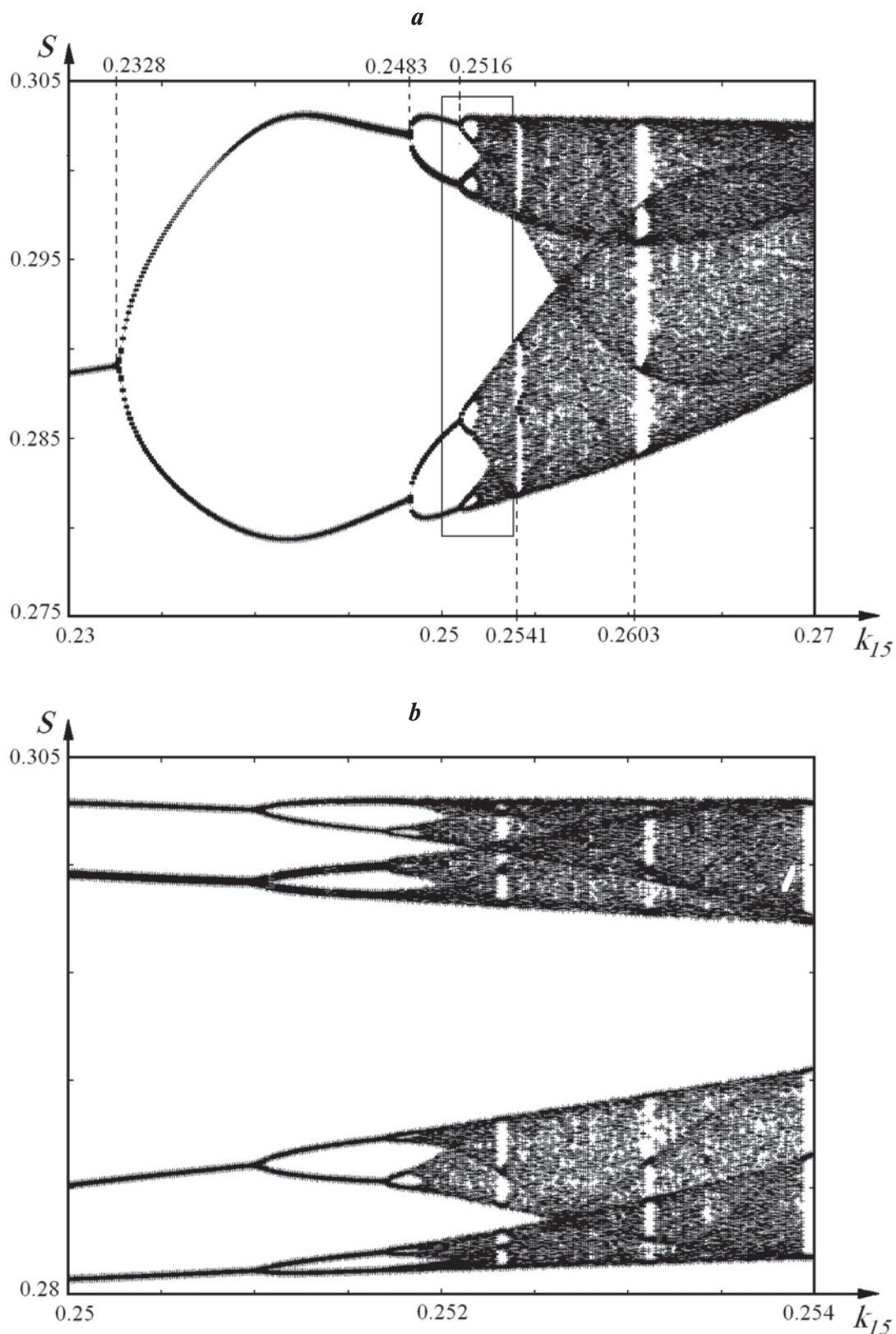


Fig. 3. Phase-parametric diagram of the system: (a) – $k_{15} \in (0.23, 0.27)$; (b) – $k_{15} \in (0.25, 0.254)$

the doubling of the period of autooscillations by the Feigenbaum scenario [36–38] at the increase in the dissipation coefficient α . This means that, for the given unstable modes of the physical system, any arising fluctuation can lead to a chaotic cyclic mode.

At $k_{15} = 0.2541$ and $k_{15} = 0.2603$ (Fig. 3, *a*), the windows of periodicity appear. The deterministic chaos is destroyed, and periodic and quasi-periodic modes are established. Outside these windows, chaotic modes hold. The identical windows of periodicity are observed also on less scales of the diagram (Fig. 3, *b*). In other words, the phase-parametric diagrams on a small scale (Fig. 3, *b*) and on a large one (Fig. 3, *a*) are analogous. This fact indicates the fractal nature of the obtained cascade of bifurcations.

As examples of the successive period doubling for the autoperiodic modes of the system by the Feigenbaum scenario, we present the projections of the phase portraits of appropriate regular attractors in Fig. 4, *a–d*. In Fig. 4, *e*, we show a regular attractor of the 6-fold quasi-periodic mode $\approx 6 \cdot 2^0$ arising in the window of periodicity at $k_{15} = 0.2543$ (Fig. 3, *a*). Fig. 4, *f* represents strange attractor $2 \cdot 2^x$ ($k_{15} = 0.267$).

The kinetics of changes in some components of the mode with strange attractor $2 \cdot 2^x$ ($k_{15} = 0.27$) is shown in Fig. 5. As compared with Fig. 2, the synchronism of oscillations of citrate S_3 and oxal-

acetate S_8 at the input and the output of the Krebs cycle is violated. The cycle stops to be strictly periodic (Fig. 2). The periodicity and the amplitudes change and become chaotic (Fig. 5). This occurs due to the violation of the synchronism of metabolic processes in the system of transport of electrons ($\text{NADH} \leftrightarrow \text{NAD}^+$) of the redox process in the respiratory chain ($Q \leftrightarrow q$) and the substrate-linked phosphorylation $\text{ADP} \rightarrow \text{ATP}$.

In Fig. 6, *a, b*, we show the projections of the phase portraits of the given strange attractor in the three-dimensional coordinate systems. The strange attractors obtained in this case are formed as a result of the folding. An element of the phase volume of such an attractor is stretched in some directions and shrinks in other ones, by conserving its stability. Therefore, we observe the mixing of trajectories in narrow places of the phase space of the system, i.e., a deterministic chaos arises.

The whole Krebs cycle consists of 8 successive reactions. In its nodal points, there occurs the transfer of electrons through NADH onto the respiratory chain. An increase in the parameter k_{15} causes an increase in the rate of the process $\text{NADH} \leftrightarrow \text{NAD}^+$ on various parts of the cycle. This causes a change in the levels of relevant metabolites on the given part. Their recurrence period in the Krebs cycle is doubled by preserving a cyclicity by the Feigenbaum scenario. In addition, the rate of substrate-linked phosphorylation $\text{ADP} \rightarrow \text{ATP}$ in

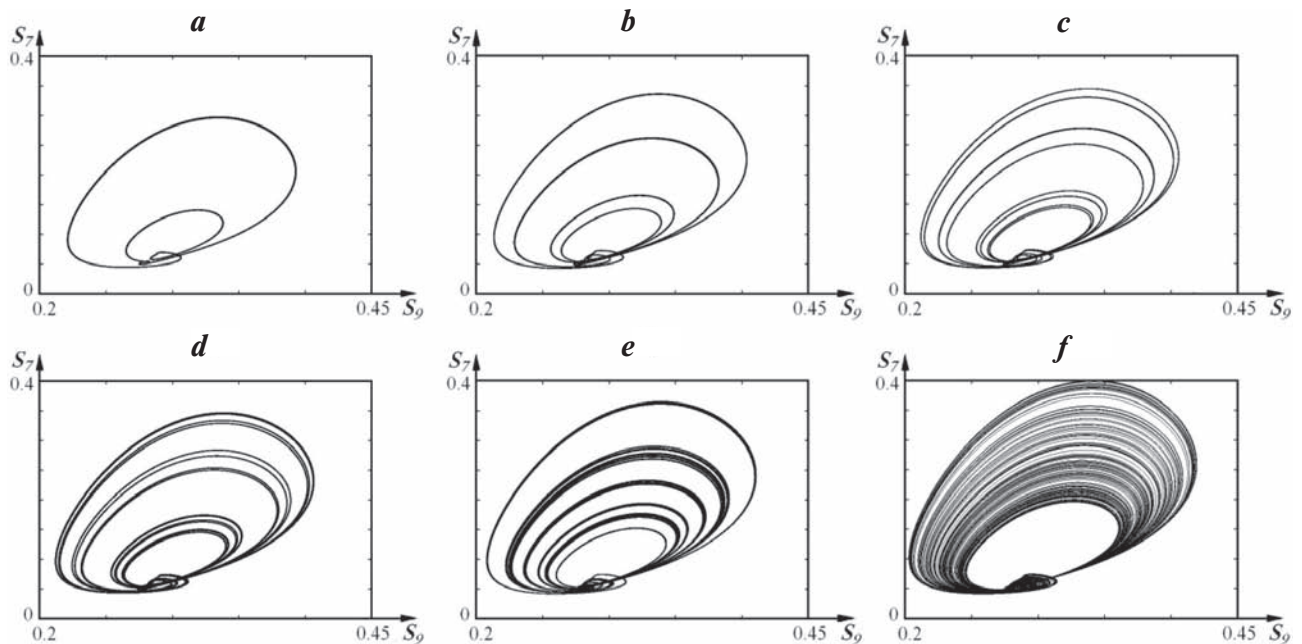


Fig. 4. Projections of the phase portraits of attractors of the system: (a) – $2 \cdot 2^0$ ($k_{15} = 0.2482$); (b) – $4 \cdot 2^0$ ($k_{15} = 0.251$); (c) – $8 \cdot 2^0$ ($k_{15} = 0.2517$); (d) – $16 \cdot 2^0$ ($k_{15} = 0.2518$); (e) – $\approx 6 \cdot 2^0$ ($k_{15} = 0.2543$); (f) – $2 \cdot 2^x$ ($k_{15} = 0.267$)

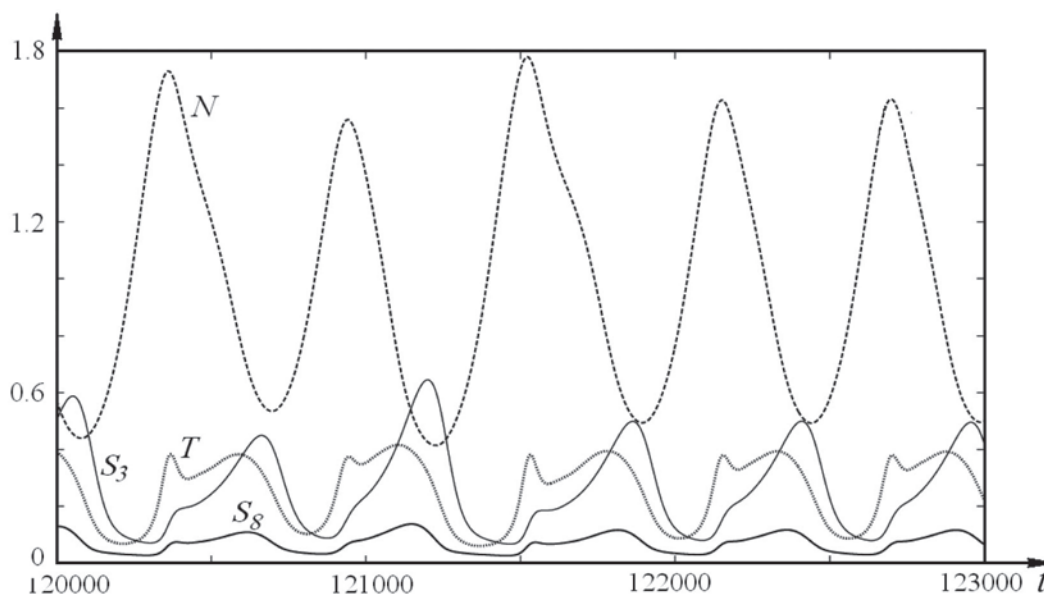


Fig. 5. Kinetic curves of components of the Krebs cycle S_3 , S_8 , N , and T in the mode of strange attractor $2\cdot 2^x$ at $k_{15} = 0.27$ and $S_8^0 = 0.001$

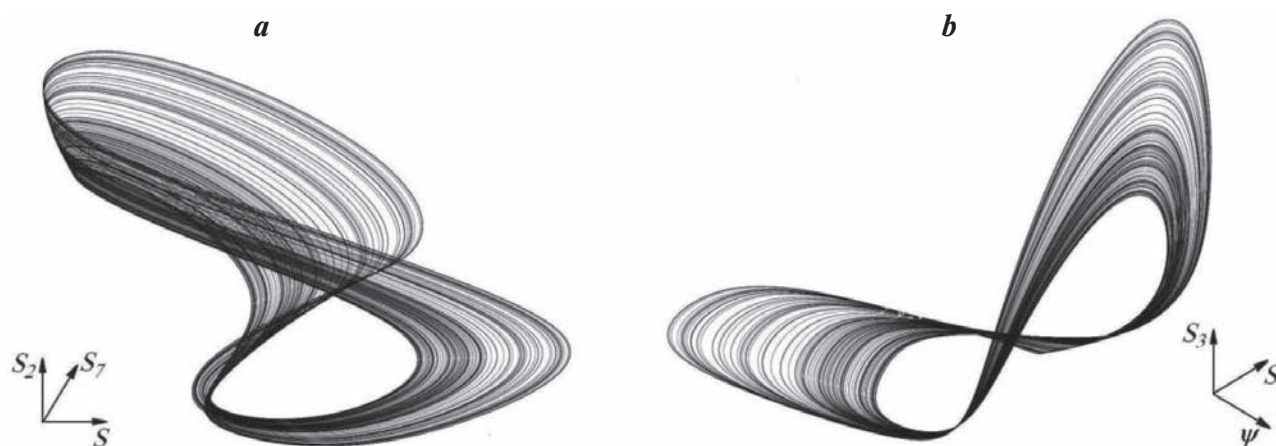


Fig. 6. Projections of the phase portraits of strange attractor $2\cdot 2^x$ at $k_{15} = 0.27$ in the three-dimensional coordinate systems: (S, S_7, S_2) (a) and (ψ, S, S_3) (b)

the respiratory chain is also increased. The arisen additional peaks on the plot of ATP correspond to those on the plot of NAD^+ (Fig. 5). Since ADP takes participation in the Krebs cycle, this causes nonlinear variations of the periodic kinetics of metabolites. The cyclicity of restoration of the level of oxalacetate S_8 is violated, which affects the subsequent cycles. In this case, a chaotic mode arises. Thus, there occurs the permanent adaptation of the metabolic process of a cell to variations in the environment. This leads to the adaptation of the whole metabolic process of a cell and to its self-organization for the conservation of homeostasis.

The results obtained allow one to estimate the stability of the metabolic process of a cell under the action of external factors and to forecast the consequence of this action on the vital activity of live organisms. The developed mathematical model can be useful for the creation of new drugs and in the study of the toxicity of various substances.

The work is supported by the project N 0113U001093 of the National Academy of Sciences of Ukraine.

САМООРГАНІЗАЦІЯ ТА ФРАКТАЛЬНІСТЬ В МЕТАБОЛІЧНИХ ПРОЦЕСАХ ЦИКЛУ КРЕБСА

В. Й. Грицай^{1,}, І. В. Мусатенко²*

¹Інститут теоретичної фізики
ім. М. М. Боголюбова НАН України, Київ;
e-mail: vgrysay@bitp.kiev.ua;

²Київський національний університет
імені Тараса Шевченка, Україна;
e-mail: ivmusatenko@gmail.com

У роботі за допомогою математичної моделі досліджується метаболізм у циклі Кребса. Знайдено автокаталітичні процеси, внаслідок яких відбувається самоорганізація в циклі Кребса і виникає циклічність в його динаміці. Досліджено структурно-функціональні зв'язки, які створюють синхронність автоперіодичного функціонування під час переносу електронів у дихальному ланцюзі та за окислювального фосфорилування. Досліджено умови порушення синхронізації процесів, збільшення кратності циклічності та виникнення хаотичних режимів. Одержана фазо-параметрична діаграма каскаду біфуркацій, яка відображає перехід до хаотичного режиму згідно зі сценарієм Фейгенбаума. Показано фрактальну природу встановленого каскаду біфуркацій. Знайдено дивні аттрактори, що утворюються завдяки формуванню складки. Одержані результати дають уявлення про структурно-функціональні зв'язки, завдяки яким відбувається самоорганізація в метаболічних процесах клітини. Побудована математична модель може бути використана для дослідження токсичного та алергічного ефектів ліків і різних речовин на метаболізм клітини.

Ключові слова: цикл Кребса, метаболічні процеси, самоорганізація, фрактальність, дивний аттрактор, сценарій Фейгенбаума.

САМООРГАНИЗАЦИЯ И ФРАКТАЛЬНОСТЬ В МЕТАБОЛИЧЕСКИХ ПРОЦЕССАХ ЦИКЛА КРЕБСА

В. И. Грицай^{1,}, И. В. Мусатенко²*

¹Институт теоретической физики
им. Н. Н. Боголюбова НАН Украины, Киев;
e-mail: vgrysay@bitp.kiev.ua;

²Киевский национальный университет
имени Тараса Шевченко, Украина;
e-mail: ivmusatenko@gmail.com

В работе при помощи математической модели исследуется метаболізм в цикле Кребса. Найденны автокаталитические процессы, вследствие которых происходит самоорганизация в цикле Кребса и возникает цикличность в его динамике. Исследованы структурно-функциональные связи, создающие синхронность автопериодического функционирования при переносе электронов в дыхательной цепи и при окислительном фосфорилировании. Исследованы условия нарушения синхронизации процессов, увеличения кратности цикличности и возникновения хаотических режимов. Получена фазопараметрическая диаграмма каскада бифуркаций, отражающая переход к хаотическому режиму согласно сценарию Фейгенбаума. Показана фрактальная природа установленного каскада бифуркаций. Найденны странные аттракторы, образуемые в результате формирования складки. Полученные результаты дают представление о структурно-функциональных связях, благодаря которым происходит самоорганизация в метаболизме клетки. Построенная математическая модель может быть использована для исследований токсического и аллергического эффектов лекарств и различных веществ на метаболизм клетки.

Ключевые слова: цикл Кребса, метаболіческие процессы, самоорганизация, фрактальность, странный аттрактор, сценарий Фейгенбаума.

1. *Krebs H. A., Johnson W. A.* // *Enzymologia.* – 1937. – N 4. – P. 148–156.
2. *Bohnensack R., Sel'kov E. E.* // *Stud. Biophys.* – 1977. – **66.** – P. 47–63.
3. *Lyubarev A. E., Kurganov B. I.* // *Molekul. Biolog.* – 1987. – **21**, N 5. – P. 1286–1296.
4. *Kondrashova M. N.* // *Biofiz.* – 1989. – **34**, Iss. 3. – P.450–457.
5. *El-Mansi E. M. T., Dawson G. C., Bryce C. F. A.* // *Comput. Applic. Biosci.* – 1994. – **10**, N 3. – P. 295–299.
6. *Ramakrishna R., Edwards J. S., McCulloch A., Palsson B. O.* // *Am. J. Physiol. Regul. Integr. Comp. Physiol.* – 2001. – **280**, N 3. – P. R695–R704.
7. *Cortassa S., Aon M. A., Marban E. et al.* // *Biophys. J.* – 2003. – **84.** – P. 2734–2755.
8. *Yugi K., Tomita M.* // *Bioinformatics.* – 2004. – **20.** – P. 1795–1796.
9. *Singh V. K., Ghosh I.* // *Theor. Biol. Med. Model.* – 2006. – **3.** – P. 27.
10. *Mogilevskaya E., Demin O., Goryanin I.* // *J. Biol. Phys.* – 2006. – **32**(3-4), P. 245–271.
11. *Gachok V. P.* *Kinetics of Biochemical Processes.* – Kiev: Nauk. Dumka, 1988. – 219 p. (in Russian).
12. *Gachok V. P.* *Strange Attractors in Biosystems.* – Kiev: Nauk. Dumka, 1989. – 237 p. (in Russian).
13. *Armiger W. B., Moreira A. R., Phillips J. A., Humphrey A. E.* *Utilization of cellulose materials in unconventional food production.* – New York: Plenum Press, 1979. – P. 111–117.
14. *Monod J.* *Recherches sur la Croissance des Cultures Bacteriennes.* – Paris: Hermann, 1942.
15. *Podgorskij V. S.* *Physiology and Metabolism of Methanol-Assimilating Yeast.* – Kiev: Naukova Dumka, 1982. (in Russian).
16. *Drozdov-Tikhomirov L. N., Rakhimova N. T.* // *Mikrobiologiya.* – 1986. – **55**, N 5. – P. 775–780.
17. *Riznichenko G. Yu.* *Mathematical Models in Biophysics and Ecology.* – Moscow, Izhevsk: Inst. of Computer. Studies, 2003. (in Russian).
18. *Watteeuw C. M., Armiger W. B., Ristroph D. L., Humphrey A. E.* // *Biotechnol. Bioeng.* – 1979. – **21**, N 5. – P. 1221–1237.
19. *Selkov E. E.* // *Europ. J. Biochem.* – 1968. – **4.** – P. 79–86.
20. *Hess B., Boiteux A.* // *Annu Rev. Biochem.* – 1971. – **40.** – P. 237–258.
21. *Goldbeter A., Lefer R.* // *Biophys J.* – 1972. – **12.** – P. 1302–1315.
22. *Goldbeter A., Caplan R.* // *Annu Rev. Biophys. Bioeng.* – 1976. – **5.** – P. 449–476.
23. *Gachok V. P., Grytsay V. I.* // *Dokl. AN SSSR.* – 1985. – **282**, N 1. – P. 51–53.
24. *Gachok V. P., Grytsay V. I., Arinbasarova A. Yu. et al.* // *Biotechnol. Bioeng.* – 1989. – **33.** – P. 661–667.
25. *Gachok V. P., Grytsay V. I., Arinbasarova A. Yu. et al.* // *Biotechnol. Bioeng.* – 1989. – **33.** – P. 668–680.
26. *Chaos in Chemical and Biochemical System / Ed. By R. Field, L. Györgyi.* – Singapore: World Scientific, 1993.
27. *Kordium V., Irodov D., Maslova O. et al.* // *Biopolym. Cell.* – 2011. – **27**, N 6. – P. 480–498.
28. *Kuznetsov S. P.* *Dynamical Chaos.* – Moscow: Nauka, 2001. (in Russian).
29. *Anishchenko V., Astakhov V., Neiman A. et al.* *Nonlinear Dynamics of Chaotic and Stochastic System. Tutorial and Modern Developments* – Berlin: Springer, 2007.
30. *Grytsay V. I.* // *Dopov. NAN Ukr.* – 2000. – N 2. – P. 175–179.
31. *Grytsay V. I.* // *Ukr. J. of Phys.* – 2001. – **46.** – P. 124–127.
32. *Grytsay V. I., Andreev V. V.* // *Matem. Modelir.* – 2006. – **18**, N 12. – P. 88–94.
33. *Grytsay V. I.* // *Romanian J. Biophys.* – 2007. – **17**, N 1. – P. 55–62.
34. *Grytsay V. I.* // *Biofiz. Visn.* – 2007. – **2**(19). – P. 92–97.
35. *Grytsay V. I., Musatenko I. V.* // *Ukr. J. Phys.* – 2013. – **58**, N 7. – P. 677–686.
36. *Feigenbaum M. J.* // *J. Stat. Phys.* – 1978. – **19**, N 1. – P. 25–52.
37. *Feigenbaum M. J.* // *J. Stat. Phys.* – 1979. – **21**, N 6. – P. 669–706.
38. *Feigenbaum M. J.* // *Comm. Math. Phys.* – 1980. – **77**, N 1. – P. 65–86.

Received 29.05.2013