Role of Krebs Cycle in the Mechanism of Stability Internal Medium and Internal Energy in an Organism in Norm and in Mechanisms of Cancer Pathology

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Abstract

The mechanisms stability of both an open non equilibrium nonlinear thermodynamic system of a human organism and the open non equilibrium nonlinear thermodynamic system of an organism’s cells were considered via study the mechanism operation of Krebs tricarboxylic acid or citric acid cycle. Just study mechanism operation of Krebs tricarboxylic acid cycle in open thermodynamic systems of both an organism and an organism’s cells gives possibility to describe the biochemical and biophysical mechanisms interactions between anabolic processes and catabolic processes considering mutual influences between anaerobic catabolic processes and aerobic catabolic processes. Also there was explained the role of Krebs cycle in biochemical and biophysical mechanisms of stability open non equilibrium nonlinear thermodynamic system of a human organism and an organism’s cells. Besides there were described the influences of Krebs cycle property on mechanisms operation of cellular capacitors contributing to maintenance stability Internal Energy both an organism and cells of an organism due to remote cellular reactions via cellular capacitors operations preceding contacts cellular reactions which leads to immune reaction on strange objects into an organism saving stability Internal Energy and Internal Medium as an organism as well as cells of an organism. Moreover there were described the mechanism of partial destruction Krebs acids cycles leading to partial violation of interactions between catabolic anaerobic processes and catabolic aerobic processes due to partial inhibition catabolic processes in Warburg effect mechanism of cancer metabolism. Also there was described the mechanisms of the offered method prevention supplementary metastasis in processes of up-to-date chemotherapy and was described practical application of this method treatment on the cancer disease patient.

Keywords: Glycolysis; Krebs tricarboxylic citric acid cycle; Anabolic endergonic processes; Catabolic aerobic exergonic processes; Catabolic anaerobic exergonic processes; Cellular capacitors; Cellular remote reactions

Introduction

The stability of both an open non equilibrium nonlinear thermodynamic system of a human organism and the open non equilibrium nonlinear thermodynamic system of an organism’s cells display balance catabolic exergonic processes and anabolic endergonic processes [1,2]. Catabolic anaerobic oxidative phosphorylation of glycolysis exerts the driving mechanism as anabolic endergonic processes as well as catabolic anaerobic exergonic processes via sharing these reverse processes in “nodal point of bifurcation anabolic and catabolic processes [NPBac]” of Acetyl-CoA [3,4] (Figure 1). TCA prolong catabolic anaerobic exergonic processes of glycolysis oxidative phosphorylation after “nodal point of bifurcation anabolic and catabolic processes [NPBac]” creating link between anaerobic catabolic exergonic processes and aerobic catabolic exergonic processes in mitochondria of cells which interact with catabolic system of Hemoglobin’s in erythrocytes of an organism’s blood. On the one hand, TCA is the link between catabolic anaerobic exergonic processes and catabolic aerobic exergonic oxidative processes. On the other hand, catabolic processes of TCA are contrary anabolic endergonic processes. Besides the influences of Krebs cycle property on mechanisms operation of cellular capacitors contribute to maintenance stability Internal Energy both an organism and cells of an organism due to remote cellular reactions via cellular capacitors operations preceding contacts cellular reactions which leads to immune reaction on strange objects into an organism saving stability Internal Energy and Internal Medium as an organism as well as cells of an organism. Just all these functions of TCA contribute to as mechanisms stability Internal Energy and Internal Medium both an organism’s cells and an organism as well as all cellular capacitors operation causing defense of Internal Energy and Internal Medium of an organism leading to stable Stationary State of an organism. Also it occurs partial destruction Krebs acids cycles leading to partial violation of interactions between catabolic anaerobic processes and catabolic aerobic processes due to partial inhibition catabolic processes in Warburg effect mechanism of cancer metabolism that cause pathologic Quasi-stationary State of an organism. Moreover there was described the mechanisms of the offered method prevention supplementary metastasis in processes of up-to-date chemotherapy and was described practical application of this method treatment on the cancer disease patient.

Krebs tricarboxylic acids cycle as the link of the mechanism maintenance stability internal energy in an organism

The ATP is produced in mitochondria via the process of oxidative phosphorylation. It occurs in such pathway: Electrons are transferred through the reducing substances of nicotinamide adenine dinucleotide (NADH) to Complex I (NADH dehydrogenase) and flavine adenine dinucleotide (FADH2) to Complex II and further through Complex III (cytochrome bc complex) then Complex IV (cytochrome c oxidase) to Complex V (ATP synthase) [5,6]. AKT is the primer for both catabolic processes and anabolic processes because AKT exerts Glycolysis advancing both catabolic exergonic and anabolic endergonic processes through “nodal point of bifurcation anabolic and catabolic processes

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The point of bifurcation anabolic and catabolic processes in metabolism of an organism.

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Krebs tricarboxylic acids cycle (TCA) in catabolic exergonic process.

![Diagram of Krebs cycle](image)

- **Glucose metabolism**
- **Fatty acids metabolism**
- **Amino acids metabolism**

**Glucose** → **Pyruvate** → **Acetyl-CoA** → **Citric acid** → **Fumarate** → **Malate** → **Oxaloacetate** → **Citric acid cycle**

**a) Transition** catabolic anaerobic processes of oxidative phosphorylation from Glycolysis to Krebs tricarboxylic acids (TCA) cycle.

**b) Glycolysis** Product "Acetyl-CoA" is as "the nodal point of bifurcation anabolic and catabolic processes [NPBac]" as well as "the one part of the link joining Glycolysis with Krebs tricarboxylic acids (TCA) cycle".

**c) Oxaloacetate** is as the closing link of circular Krebs tricarboxylic acids (TCA) cycle as well as "the second part of the link joining Glycolysis with Krebs tricarboxylic acids (TCA) cycle".

**d) The joint** between Acetyl-CoA and Oxaloacetate produces Citric acid as the primary link of driving mechanism Krebs tricarboxylic citric (TCA) cycle which governs direction reaction among Glycolysis and Krebs tricarboxylic citric (TCA) cycle.

**e) The final** Catabolic Products of Krebs tricarboxylic citric (TCA) cycle are Carbon dioxide and Hydrogen ion.

**f) Hydrogen ion** is oxidized by Oxygen resulting in Water.

**Figure 2**: Krebs tricarboxylic acids cycle (TCA) in catabolic exergonic process.

**Figure 1**: The point of bifurcation anabolic and catabolic processes in metabolism of an organism.

**a) Nodal point** of bifurcation anabolic and catabolic processes in "Nodal point of bifurcation anabolic and catabolic processes [NPBac]."

**b) Moderate** metabolic processes displaying balance anabolic and catabolic processes in able-bodied tissue.

**c) Accumulation** of energy into lactic acid for anabolic processes.

**d) Normal** excretion substances via catabolic oxidative processes in able-bodied tissue.

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An organism’s cells accept Oxygen from Oxyhaemoglobin of erythrocytes transporting iron of their haeme from Fe²⁺ into Fe³⁺ plus free oxygen, i.e., forming Haemoglobin with also releasing free oxygen.
Free oxygen adds electron, which is produced by serial transformations NAD\(^+\)→NADH and FAD\(^+\)→FADH\(_2\), and forms ROS producing moderate quantity dissolved Superoxide:

\[ \text{b) } [\text{Hb-Fe}^3+2e] \rightarrow \text{Hb-Fe}^3+ + O_2 \]  
\[ [\text{cytc-Fe}^3+ + e] \rightarrow [\text{cytc-Fe}^2+] \]
\[ \text{c) } [\text{cytc-Fe}^3+ + e] \rightarrow \text{Hb-Fe}^3+ + O_2 \]  
\[ [\text{cytc-Fe}^3+ + e] \rightarrow \text{cytc-Fe}^2+ + O_2 \]

The dissolved Superoxide is transported by electron transport system (ETS) across cellular membranes and mitochondrial membranes binding with cytochrome c that transforms iron of cytochrome c from Fe\(^3+\) to Fe\(^2+\) and form oxidized cytochrome c. Then the oxidized cytochrome c adds electron and release free Cytochrome C with free Oxygen. Oxygen react with Hydrogen ion from Krebs cycle producing Water due to operation of cytochrome c oxidase, i.e., elimination of free oxygen (Figures 2 and 3).

\[ \text{d) } \text{cytc-Fe}^3+ + O_2^- \rightarrow [\text{cytc-Fe}^3+2O_2]; \text{e) } [\text{cytc-Fe}^3+2O_2] + e \rightarrow \text{cytc-Fe}^2+ + O_2 \]
\[ \text{f) cytc-Fe}^3+ + O_2^- \rightarrow \text{cytc-Fe}^2+ + O_2 \]  
\[ \text{g) } 4\text{H}^+ + \text{O}_2 = 2\text{H}_2\text{O} \]

However, the quantity of consumed oxygen in the capillaries of lung’s alveoli are considerably more then the produced Hydrogen ion in Krebs cycle [5,6]. Therefore, there are formed considerably more quantity of surplus Superoxide (O\(_3^\cdot\)) from surplus oxygen (O\(_2\)) via adding electron:

\[ \text{h) } n[\text{O}_2] + n[e^-] \rightarrow n[\text{O}_3^\cdot] \]

Then surplus Superoxide (O\(_3^\cdot\)) is transported by electron transport system (ETS) into mitochondria. Further surplus Superoxide (O\(_3^\cdot\)) reduces Ferric iron [Fe\(^3+\)] into Ferrous iron [Fe\(^2+\)] with oxygen:

\[ \text{i) } O_3^\cdot + \text{cytc-Fe}^3+ \rightarrow \text{cytc-Fe}^2+ + O_2 \]

Simultaneously Reactive Oxygen Species (ROS) is generated by NOX (NADPH oxidase) and Duox due to activity mitochondrial aerobic catabolic processes forming of this surplus quantity of mitochondrial Superoxide (O\(_3^\cdot\)) which don’t continue processes of anaerobic oxidative phosphorylation and don’t lead down to final products CO\(_2\) and H\(_2\)O. Just NAD\(^+\) and FAD are transformed: NAD\(^+\) is transformed into reduced molecule NADH due to NADPH dehydrogenase operation, and FAD is transformed into reduced molecule FADH\(_2\). Then NADH is back oxidized into NAD\(^+\) by NADH oxidation at Complex I, and pair of electrons from NADH is released passing via series of electron transport carriers of electron transport system (ETS) to coenzyme Q into complex I and further through Complex III (cytochrome bc complex) then Complex IV (cytochrome c oxidase) to Complex V (ATP synthase) into the intermembrane space [5,6].

The partial supplementary Superoxide anion [O\(_3^\cdot\)] is accumulated into forming Reactive Oxygen Species [ROS], and the rest supplementary Superoxide anion [O\(_3^\cdot\)] reacts with hydrogen cations forming hydrogen peroxide and oxygen:

\[ \text{j) } 2O_3^\cdot + 2\text{H}^+ \rightarrow \text{H}_2\text{O}_2 + O_2 \]

Further there is happened Haber –Weiss reaction of iron catalysed by superoxide transformations being passed into Fenton reaction which is applied to mitochondria producing free radicals [‘OH] in such mode [5,6]:

\[ \text{k) } \text{cytc-Fe}^3+ + \text{H}_2\text{O}_2 \rightarrow \text{cytc-Fe}^3+ + \text{OH} + \text{‘OH} \]
\[ \text{cytc-Fe}^3+ + O_2^- \rightarrow \text{cytc-Fe}^2+ + O_2 \]
\[ O_3^\cdot + \text{H}_2\text{O}_2 \rightarrow \text{OH} + \text{‘OH} + O_2 + \text{Fe}^{3+} \]

Thus these converting’s of supplementary respired oxygen lead to creation complex ROS/H\(_2\)O/ free radicals (‘OH) which pass through mitochondrial membranes, through cytoplasm and nuclear membranes penetrating into nucleus to nuclear DNA. On the one hand, ROS is neutralized by glutathione peroxidise (GPX) and phospholipid hydroperoxide glutathione peroxidise (PHGPX) in normal G1/S phases of cellular cycle [5,6]. On the other hand, moderate anabolic processes transit into intensive anabolic processes in nucleus, and simultaneously moderate catabolic processes transits into intensive catabolic processes in mitochondria mechanism maintenance stability chemical potential of cytoplasmin (β\(_{\text{m,mito}}\)) and maintaining balance catabolic processes and anabolic processes in cytoplasmin that leads to production surplus complex ROS/H\(_2\)O in normal G\(_2\) phase of cellular cycle [6]. Complex ROS/H\(_2\)O generates superoxide (O\(_3^\cdot\)) into nucleus which induces free radicals (‘OH). Free radicals (‘OH) react with nuclear DNA (nDNA) and induce process replication via realizing of 2nDNA [6-13].

\[ \text{l) } \text{‘OH} + \text{H}_2\text{nDNA-DNA} \rightarrow \text{H}_2\text{O} + \text{H}^+\text{nDNA-DNA} \]
\[ \text{O}^+ + 2\text{H}_2\text{O} \rightarrow 2\text{H}^+ + 2\text{OH}^- \]
\[ 2\text{H}^+\text{nDNA-DNA} + 2\text{H}^+ \rightarrow 2\text{DNA-H}^+ + 2\text{DNA-H}^- \]
\[ 2\text{DNA-H}^+ + 2\text{OH}^- \rightarrow 2\text{DNA} + \text{H}_2\text{O} \]

Thus the free radicals (‘OH and H\(^+\)) are neutralized in final G\(_2\) phase of DNA replication. Then it occurs M phase of cellular cycle of Mitosis in cell division that transfers the new cells into G\(_1\) phase of normal cellular cycle. Thus, nuclei DNA (nDNA) of formed new cells are not subjected to ruining capability of ROS/H\(_2\)O/ free radicals in normal development cellular cycle [2,5,6]. Such mutual influences between nucleus and mitochondria induce chemical potentials of cellular cytoplasmin’s in G\(_1\), G\(_S\)/G\(_1\), and M/G\(_2\) phases normal cellular cycle which are related to chemical potentials an organism maintaining stable Internal Energy and Internal Medium both in an organism and cells of an organism via cellular capacitors, nuclear capacitors and mitochondrial capacitors operations in common interconnection and mutual interdependence [2,4-6]. Just these related chemical potentials are supported by the operations of nuclear capacitors, mitochondrial capacitors and cellular capacitors via generating relative resonance waves between cells and an organism in norm [4-6,12]. Just Krebs tricarboxylic acids cycle carries out the role of main driving mechanism of production supplementary superoxide (O\(_3^\cdot\)): a) The quantity Hydrogen ion (H\(^+\)) produced in TCA, reacts with free Oxygen (O\(_2\)) and forms Water (H\(_2\)O) that eliminates free Oxygen in liquids of an organism and cells. b) The supplementary oxygen (O\(_3^\cdot\)), which did not react with Hydrogen ion (H\(^+\)) produces supplementary superoxide (O\(_3^\cdot\)) which generates free radicals exerting replications in G\(_1\)/S, G\(_1\), M/G\(_2\) phases of normal cellular cycle [6]. On the one hand, TCA influences on anabolic processes due to production supplemental superoxide (O\(_3^\cdot\)) and generating Free radicals (‘OH) which induce nuclear proliferative processes in G\(_2\) phase of cellular cycle [2-6]. On the other hand, the pathway of Glycolysis is divided in “nodal point of bifurcation of anabolic and catabolic processes” [NPBac] of Acetyl-CoA, in which it occurs mutual influences between anabolic endergonic processes and catabolic exergonic processes [3,4] (Figure 1). Just Krebs tricarboxylic citric acid cycle, joining catabolic anaerobic and aerobic processes, is the link which influences on maintenance stability of balance catabolic exergonic processes and anabolic endergonic processes through its main crucial point of molecules citric acids and “nodal point of bifurcation of anabolic and catabolic processes [NPBac] in Acetyl-CoA”. Thus it occurs mutual influences between anabolic endergonic processes and Krebs tricarboxylic acid (TCA) cycle through crucial molecules of Acetyl-CoA and Citric Acids for maintenance stability Internal Energy as in an organism as well as in cells of an organism (Figures 2 and 3). Besides interactions between
stable Krebs tricarboxylic acid cycle and advancing anabolic processes induce different charges on Internal and External Membranes of cellular capacitors, nuclear capacitors, mitochondrial capacitors and the other organelles' capacitors that contributes to maintenance stability Internal Energy of all cells via operations relative resonance waives between them [4,12]. Thus, TCA cycle are the joint mechanism which exerts as the mechanisms of biochemical maintenance stability Internal Energy and Internal Medium of an organism [9,10] as well as the mechanism of biophysical maintenance stability Internal Energy and Internal Medium of an organism [4,12] (Figures 2 and 3).

The role of Krebs tricarboxylic acids cycle in cancer metabolism

Cancer metabolism is the result of oncogenes operation as the etiologic factors [14-17]. Just there were described following oncogenes manifestations: The operations kinase corresponded to influences of genes on mutation processes [18] Rb1 is phosphorylated by cyclin-dependent kinases (CDKs) and plays important role in DNA replication and in cells division in G2/M cellular phases, and also Rb1, is subjected to mutation in some tumors [18]. It occurs the suppression of p53 expression by RNA, in some tumors [18]. Intruding into cellular genome of v-oncogene, the cells were subjected to accept of accelerated cellular rhythm of this v-oncogene into cellular Genome exerting huge anabolic endergonic processes in cancer tissue /+2000H+/ that leads to shift balance catabolic exergonic processes and anabolic endergonic processes into excessive anabolic endergonic processes of cancer metabolism, as compared with normal balance catabolic exergonic processes and anabolic endergonic processes in able-bodied tissues/2+H+ [2,3] (Figure 4). The excessive anabolic endergonic processes in cancer metabolism take up huge quantity energy and Acetyl-CoA that cause overloading of “nodal point of bifurcation anabolic and catabolic processes” [NPBag] remaining lack of energy and Acetyl–CoA for catabolic oxidative processes promoting cancer cells’ survival via showing Apoptosis Resistance. Just lack Acetyl-CoA causes the partial suppression oxidative phosphorylation of catabolic exergonic processes, i.e., Krebs citric acids cycle, in cancer metabolism [3]. The increase of lactic acids production in cancer metabolism is the necessary endergonic mechanism accumulation energy for huge anabolic processes in condition glycosis metabolism and enormous consumption of energy in cancer metabolism [18]. Just partial suppression of Krebs tricarboxylic acids cycle occurs in Oxaloacetate link of circular metabolic pathway between these Krebs tricarboxilic citric (TCA) cycles. [mitochondrialTCA, cytoplasmic TCA and an organism’s extracellular TCA] are making circular pathways between these Krebs tricarboxilic citric (TCA) cycles. a) Oxaloacetates join three Krebs tricarboxilic citric (TCA) cycles [mitochondrialTCA, cytoplasmic TCA and an organism’s extracellular TCA] making circular pathways between these Krebs tricarboxilic citric (TCA) cycles. b) The final Products of all three Krebs tricarboxilic citric (TCA) cycles [mitochondrialTCA, cytoplasmic TCA and an organism’s extracellular TCA] are Carbon dioxide ions and Hydrogen ions. c) The final Catabolic Products of Carbon dioxide ions are transferred to lung’s alveoles by Carboxyhaemoglobin in blood of an organism where Carbon dioxide ions turn into Carbon dioxide. d) Oxygen ions turn into Superoxide which reacts with Hydrogen ions resulting in Oxygen ion and Water in lung’s alveoles.

The increase of lactic acids production in cancer metabolism is the result of oncogenes operation as the etiologic factors [14-17]. Just there were described following oncogenes manifestations: The operations kinase corresponded to influences of genes on mutation processes [18] Rb1 is phosphorylated by cyclin-dependent kinases (CDKs) and plays important role in DNA replication and in cells division in G2/M cellular phases, and also Rb1, is subjected to mutation in some tumors [18]. It occurs the suppression of p53 expression by RNA, in some tumors [18]. Intruding into cellular genome of v-oncogene, the cells were subjected to accept of accelerated cellular rhythm of this v-oncogene into cellular Genome exerting huge anabolic endergonic processes in cancer tissue /+2000H+/ that leads to shift balance catabolic exergonic processes and anabolic endergonic processes into excessive anabolic endergonic processes of cancer metabolism, as compared with normal balance catabolic exergonic processes and anabolic endergonic processes in able-bodied tissues/2+H+ [2,3] (Figure 4). The excessive anabolic endergonic processes in cancer metabolism take up huge quantity energy and Acetyl-CoA that cause overloading of “nodal point of bifurcation anabolic and catabolic processes” [NPBag] remaining lack of energy and Acetyl–CoA for catabolic oxidative processes promoting cancer cells’ survival via showing Apoptosis Resistance. Just lack Acetyl-CoA causes the partial suppression oxidative phosphorylation of catabolic exergonic processes, i.e., Krebs citric acids cycle, in cancer metabolism [3]. The increase of lactic acids production in cancer metabolism is the necessary endergonic mechanism accumulation energy for huge anabolic processes in condition glycosis metabolism and enormous consumption of energy in cancer metabolism [18]. Just partial suppression of Krebs tricarboxylic acids cycle occurs in Oxaloacetate link of circular metabolic pathway between these Krebs tricarboxilic citric (TCA) cycles. [mitochondrialTCA, cytoplasmic TCA and an organism’s extracellular TCA] are making circular pathways between these Krebs tricarboxilic citric (TCA) cycles. a) Oxaloacetates join three Krebs tricarboxilic citric (TCA) cycles [mitochondrialTCA, cytoplasmic TCA and an organism’s extracellular TCA] making circular pathways between these Krebs tricarboxilic citric (TCA) cycles. b) The final Products of all three Krebs tricarboxilic citric (TCA) cycles [mitochondrialTCA, cytoplasmic TCA and an organism’s extracellular TCA] are Carbon dioxide ions and Hydrogen ions. c) The final Catabolic Products of Carbon dioxide ions are transferred to lung’s alveoles by Carboxyhaemoglobin in blood of an organism where Carbon dioxide ions turn into Carbon dioxide. d) Oxygen ions turn into Superoxide which reacts with Hydrogen ions resulting in Oxygen ion and Water in lung’s alveoles.

e) The final Catabolic Products of Hydrogen ions react with Oxygen ion resulting Water due to Cytochrome C oxidase operation in mitochondrial Cytochrome C. Figure 3: Interactions between anaerobic catabolic processes and aerobic catabolic processes via Krebs.

f) O3 + e → O2*

Then the great quantity of surplus Superoxide (O3*) considerably more than in normal tissue, is transported by electron transport system (ETS) into mitochondria in cancer tissue. Further the great quantity of surplus Superoxide [O3] reduces Ferric iron [Fe3+] into Ferrous iron [Fe2+] realising also oxygen:

g) O3*+cytc-Fe3+→ cytc-Fe2+O2. Simultaneously surplus Reactive Oxygen Species (ROS) is generated by NOX (NADPH oxidase) and Duox due to activity mitochondrial aerobic catabolic processes forming of this surplus quantity of mitochondrial superoxide [O3*].

NAD+ and FAD are transformed in processes of surplus Superoxide [O3] formation in cancer tissue metabolism in more intensity than in normal tissue metabolism: NAD+ is transformed into reduced molecule NADH due to NADPH dehydrogenase operation, and FAD is transformed into reduced molecule FADH2. Then NADH is back oxidized into NAD+ by NADH oxidase operation at Complex I, and lung’s alveoli are considerably more than the produced Hydrogen ion in partial suppressed Krebs cycle in cancer tissue and ever than the produced Hydrogen ion in Krebs cycle in an able-bodied tissue [6]. Therefore, there are formed surplus Superoxide (O3*) in cancer tissue, considerably more quantity than in an able-bodied tissue, due to great quantity of surplus oxygen (O2) via adding electron in cancer tissue [6].
pair of electrons from NADH is released passing via series of electron transport carriers of electron transport system (ETS) to coenzyme Q into Complex II and further through Complex III (cytochrome bc complex) then Complex IV (cytochrome c oxidase) to Complex V (ATP synthase) into the intermembrane space [5,6]. The great quantity of Reactive Oxygen Species (ROS) is generated in cancer tissue by NOX (NADPH oxidase) and Duox due to activity mitochondrial aerobic catabolic processes forming of this great quantity of surplus quantity of mitochondrial superoxide [O$_2^-$]. Also the partial surplus Superoxide anion [O$_2^-$] is accumulated into forming Reactive Oxygen Species [ROS]. However the great quantity of Superoxide anion [O$_2^-$] is subjected to dismutation by manganese superoxide dismutase (MnSOD) and copper, zinc superoxide dismutase (Cu, ZnSOD) converting into great quantity of hydrogen peroxide in cancer tissue: 

$$2O_2^- + 2H^+ = H_2O_2 + O_2$$

Subsequently it is happened Haber – Weiss reaction of iron catalysed by superoxide transformations which is passed into Fenton reaction which is applied to mitochondria producing free radicals [OH$^-$] in such mode [5,6,19-27]

$$Fe^{3+} + O_2^- \rightarrow Fe^{2+} + O_2$$

$$2Fe^{2+} + H_2O_2 \rightarrow Fe^{3+} + OH^- + OH + O_2 + Fe^{2+}$$

The partial abundance hydrogen peroxide [H$_2$O$_2$] from ROS is detoxified by mitochondrial glutathione peroxide (GPx) and phospholipid hydroperoxide glutathione peroxide (PHGPx) in G1/S phases of cellular cycle: Glutathione (GSH) is transformed into glutathione peroxide [GPx] caused by superoxide transformations which is passed into Fenton reaction. Thus, ROS/H$_2$O$_2$ detoxifying system forms glutathione disulphide (GSSG) in mitochondrial glutathione peroxide (GSH) is transformed into oxidized glutathione which is reduced back to GSH by glutathione reductase which promote the full neutralization of ROS/H$_2$O$_2$. Also the partial surplus hydrogen peroxide (H$_2$O$_2$) from ROS is transformed into water (H$_2$O) by glutathione peroxide (GPx).

$$2H_2O_2 \rightarrow 2H_2O + O_2$$

$$H_2O_2 + GSH \rightarrow GSSG + 2H_2O$$

The majority of abundance complex ROS/H$_2$O$_2$/free radicals operates in G$_i$ phase cellular cycle, inducing process replication via realizing of 2nDNA [6,13].

$$O^2^- + H_2O_2 \rightarrow 2H^+ + 2OH^-$$

$$2H^+ + 2nDNA-DNA \rightarrow 2nDNA-H^+ + 2nDNA-H^-$$

Thus, ROS/H$_2$O$_2$/free radicals exert excessive processes of DNA replication which promote the full neutralization of ROS/H$_2$O$_2$/free radicals, eliminating their ruining properties in G$_i$ phase oncologic cellular cycle. Also the great acceleration of cellular cycle, induced by oncogene, leads to unnoticeable G$_i$ phase in oncologic cellular cycle. Division cell in M phase oncologic cellular cycle leads to forming new cells in G$_i$/S cellular cycle due to acceleration cellular cycle and unnoticeable G$_i$ phase cellular cycle. The great acceleration of cellular cycle, induced by oncogene, with combination of abundance ROS and excessive processes of DNA replication causing neutralization ROS/H$_2$O$_2$/free radicals eliminates these incompatible resisted situations in metabolism of cancer cells, induced by mechanism of abundance ROS function: On the one hand, large amount of ROS production with hydrogen peroxide in mitochondria of cancer cells which would lead to apoptotic damage of cancer cells, and, on the other hand, cancer...
metabolism is characterized by Apoptosis Resistance [3,36-38]. Just it is the mechanism Apoptosis Resistance in oncologic cellular cycle. Oncologic cellular cycle is characterized by expression huge anabolic processes in cellular oncogenesis. Hence the excessive shift of the balance anabolic and catabolic processes into abundance anabolic processes with accelerating cellular cycle in tumor tissue advances cellular cycle in cellular oncogenesis via G1/S, G2 and M/G0/S phases which create chemical potentials unrelated to chemical potentials as an organism as well as between new formed cells that is driver mechanism of proliferative processes leading to formation Warburg effect with excessive proliferative processes, irrepressible cancer growth, unhealed cancer wounds, mechanisms of metastasis and Apoptosis Resistance [3-6]. Thus, it is formed Warburg effect mechanism of "aerobic glycolysis" in cancer metabolism, versus Pasteur effect "incompatibility aerobic oxidation with glycolysis" in able-bodied tissue [3,19-26] (Figure 4 and 5). Thus, cellular oncogenesis exhibits abundance ROS which is also driving mechanism of excessive processes of DNA replication in G2 phase cellular cycle leading to excessive cancer cells proliferative processes [3,5,6].

Highlight of cancer genesis: As outcome of oncogenes operation, the huge anabolic processes cause huge consumption of energy and Acetyl-CoA and partial suppress the catabolic processes in cancer tissue. Lactic acids accumulate energy for anabolic processes in condition glycolysis metabolism remaining the part of the energy oxidative phosphorylation in Krebs tricarboxylic acids cycle which generate energy for maintenance stability Internal Energy of an organism and cells of an organism how temperature 36,6°C-37.0°C by which all enzymes operate etc. Also this energy is used for cancer cells survival displaying Apoptosis Resistance. The partial suppressed TCA produces considerably less quantity of Hydrogen ions than in norm, that leads to remained considerably more free oxygen causing productions great quantity of abundance complex ROS/H2O2/Free radicals which induce process replication via realizing of 2ndDNA exerting excessive irrepressible proliferative processes. Thus, it forms Warburg effect mechanism of "aerobic glycolysis" which creates Apoptosis Resistance of cancer cells, metastasis, irrepressible cells growth, unhealed cancer ulcer and so on.

Exertion expression Krebs tricarboxylic citric acids cycle in cancer metabolism for prevention supplementary metastasis in processes of up-to-date chemotherapy

The mechanisms of prevention supplementary metastasis in processes of up-to-date chemotherapy: There are lot of theories of cancer metastasis mechanism [3,36-38]. Some theories explain mechanism of cancer metastasis because of the low expression of K+ in the K+ channel of malignant tumors [36-38]. We have chosen the described mechanism of metastasis based on explanation mechanism of Warburg effect by Ponizovskyi MR [3] and also the role of mitochondrial frataxin protein in mechanism of cancer metastasis [3,39]. "The huge anabolic processes cause the blockade of excretion (outflow) the synthesized high-molecular substances from cancer tissue by the pathway of oxidative metabolism because of overload "NPBac" and "lack of Acetyl-CoA" for oxidative processes" in cancer tissues (Figure 4). Therefore, the alternative pathway of high-molecular substance excretion takes place within separate cells. The viable separate cells transit from the malignant tumor tissue into the Internal Medium of the organism (blood or lymph) and, being diffused by lymph or blood, get to a healthy place of extracellular matrix without overloaded "NPBac" and "lack of Acetyl-CoA" and form metastases. Just it is happened the suppression of mitochondrial frataxin due to overloaded "NPBac" and "lack of Acetyl-CoA" promoting suppression catabolic oxidative processes [3-39]. Also the overloaded "NPBac" and "lack Acetyl-CoA", which is the carrier of K+ ions in Kv channel, contribute to the low expression of K+ in the K+ channel of malignant tumors in condition of cancer metastasis [36-38]. Thus, the suppression catabolic processes due to overloaded "NPBac" and "lack of Acetyl-CoA touches on TCA cycle via inhibition link of transferring Oxaloacetates from an organism to cancer cells violating link between mitochondrial Citric Acids cycle and mitochondrial system of cytochrome due to decreased production of quantity Hydrogen ions in Krebs tricarboxylic acids cycle (Figure 5). The study of cancer metastasis mechanism contributes to use of citric acids from citric juice, in which there is preserved the enzymes for Citric Acids cycle, that exerts as expression Krebs tricarboxylic citric acids cycle in cancer cells as well as increases Acetyl-CoA eliminating overloaded "NPBac" and causing prevention additional metastasis in processes of up-to-date chemotherapy. Really excessive quantity of citric acids with appropriate enzymes increases quantity Acetyl-CoA due to maintenance stable index of Equilibrium Constant reaction in Krebs tricarboxylic acids cycle [1] Acetyl-CoA → Oxaloacetate → Citric acid. This reaction in Krebs tricarboxylic acids cycle moves right saving stable Equilibrium Constant of reaction: Acetyl-CoA + Oxaloacetate → Citric acid. However, this reaction moves left in condition of excessive increase of quantity citric acid saving stable Equilibrium Constant via increase quantity of Acetyl-CoA too: Acetyl-CoA + Oxaloacetate → Citric acid. Increase quantity of Acetyl-CoA eliminates overloaded "NPBac" that cause prevention additional metastasis in processes of up-to-date chemotherapy. Besides excessive increased quantity citric acid exerts Citric Acids cycle causing expression of catabolic exergonic processes that induce also mitochondrial frataxin protein operation in cytochrome c catabolic aerobic processes inhibiting cancer metabolism and preventing cancer metastasis in processes of up-to-date chemotherapy.

Some clinical observation of using offered method prevention cancer metastasis: At 2010 year the man Pol. was examined and was found out a polyp in the large intestine. The repeated investigation was made at December 10, 2014 and was found cancer of large intestine. There was made the operation in which proceeding it was found the multiple metastasis in visceral peritoneum. The sick Pol. has refused himself from the offered alternative therapy of "Prolonged medical starvation during 45 days with small dosage cytotoxic remedy" [40-43]. He has chosen up-to-date method of chemotherapy. At December 2014, the sick Pol. has begun to receive chemotherapy treatment with Fluouracil and Erbitux immunotherapy. Taking into account the above described mechanisms of the role citric acids operation in Citric acids cycle, I have recommended to use citric juice from the squeezed two citrons during day, i.e., over the entire circadian period, which should be diluted in 1 litre water for prevention intestine irritation from great concentration of citric acids. This method must be used simultaneously with chemotherapy, as the additional method for prevention new metastasis. The patient Pol. has drunk the citric juice prepared in such mode during two months. Then patient Pol. has drunk the citric juice from the squeezed one citron daily which was diluted in 1 litre water. The weekly examinations show that there were not found new metastasis by the patient Pol. during the period from December 2014 till September 2016. The state of the patient Pol. is satisfactory.

Conclusions

The common pathway of both Glycolysis and Krebs tricarboxylic acid cycle gives possibility to estimate as combination related catabolic exergonic anaerobic processes and catabolic exergonic aerobic processes as well as their interactions with opposed them anabolic endergonic processes. The common pathway of both Glycolysis and
Krebs tricarboxylic acid cycle is the basic link of balance catalytic exergonic processes and anabolic endergonic processes which is also the basic link of mechanism maintenance stability Internal Energy of an organism and cells of an organism according first law of thermodynamics. Acetyl-CoA is the joining link in "nodal point of bifurcation catalytic and anabolic processes [NPBac]" between anabolic endergonic processes and catalytic exergonic anerobic processes exhibiting oxidative phosphorylation processes of Glycolysis and Krebs tricarboxylic cycle. Oxaloacetates are the joining links between Krebs tricarboxylic cycles of an organism’s cells and Krebs tricarboxylic cycles of an organism making the joint between catalytic exergonic anaerobic processes and catalytic exergonic aerobic exergonic oxidative processes. Citric acid is the primary link of driving mechanism Krebs tricarboxylic citric cycle which governs direction reaction among Glycolysis and Krebs tricarboxylic citric cycle as the product of reaction between Acetyl-CoA and Oxaloacetate. It was described mechanism maintenance stability Internal Energy of an organism in processes of respiratory in norm. The accelerated cellular rhythm of the v-oncogenes intrusted into cellular Genome exerts accelerated proliferative processes with huge anabolic endergonic processes of cancer oncogenesis leading to shift balance catalytic exergonic processes and anabolic endergonic processes into excessive anabolic endergonic processes of cancer metabolism. Consuming huge quantity both energy and Acetyl-CoA for excessive anabolic endergonic processes, it occurs overload of “nodal point of bifurcation anabolic and catalytic processes” [NPBac] in point of Acetyl-CoA that leads to remaining lack of energy and Acetyl–CoA for catalytic oxidative processes and causes the partial suppression oxidative phosphorylation of catalytic exergonic processes in cancer metabolism forming mechanism of Warburg effect. Lactic acids accumulate energy for excessive anabolic processes in condition glycolysis metabolism remaining the part of the energy oxidative phosphorylation in Krebs tricarboxylic acids cycle for cancer cells’ survival exhibiting Apoptosis Resistance of cancer metabolism. The partial suppression of Krebs tricarboxylic acids cycle occurs in Oxaloacetates link of its circular pathway between an organism and mitochondria that separates Citric Acids Cycle of an organism from Citric Acids Cycle of cancer cells, making autonomic mechanism of cancer cells development. The partial suppressed Krebs tricarboxylic acids cycle produces considerably less quantity of Hydrogen ions than in norm, that leads to remained considerably more free oxygen causing productions great quantity of Superoxide (O$_2^-$), considerably more than in norm. Superoxide (O$_2^-$) binds with cytochrome c that transforms iron of cytochrome c from Fe$^{2+}$ into Fe$^{3+}$ and form oxidized cytochrome c. The oxidized cytochrome c adds electron and release free Cytochrome C with free Oxygen which reacts with Hydrogen ion from Krebs tricarboxylic acids cycle resulting in production Water. The quantities of consumed oxygen in the capillaries of lung’s alveoli are considerably more than the produced Hydrogen ion in Krebs tricarboxylic acids (TCA) cycle that leads to form surplus Superoxide (O$_2^-$) which reduces Ferric iron [Fe$^{3+}$] into Ferrous iron [Fe$^{2+}$] with production oxygen, and this great quantity surplus Superoxide (O$_2^-$) are subjected to dismutation into great quantity of hydrogen peroxide in cancer tissue. The partial surplus Superoxide anion (O$_2^-$) is accumulated into forming Reactive Oxygen Species [ROS]. Haber-Weiss reaction of iron catalysed by superoxide transformations which is passed into Fenton reaction which is applied to mitochondria producing free radicals [‘OH]. The abundance complex ROS/H$_2$O$_2$/free radicals operates in G, phase cellular cycle of cancer cells and induce process accelerating replication via realizing of 2nDNA leading to irrepressible proliferation of cancer cells. The use of citric acids in citric juice, in which it was preserved the enzymes for Citric Acids cycle, exerts as expression Krebs tricarboxylic citric acids cycle in cancer cells as well as increases Acetyl-CoA eliminating overloaded “NPBac” and causing prevention additional metastasis in processes of up-to-date chemotherapy. There was described the some preliminary clinical positive results of use the offered method of prevention additional metastasis with chemotherapy by cancer patient.

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This article is dedicated to the memory of my daughter TM Ponizovskyi.

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