Mitochondrial Dysfunction in Depression

Yashika Bansal and Anurag Kuhad*

Pharmacology Research Laboratory, University Institute of Pharmaceutical Sciences, UGC-Centre of Advanced Study, Panjab University, Chandigarh-160 014 India

Abstract: Background: Depression is the most debilitating neuropsychiatric disorder with significant impact on socio-occupational and well being of individual. The exact pathophysiology of depression is still enigmatic though various theories have been put forwarded. There are evidences showing that mitochondrial dysfunction in various brain regions is associated with depression. Recent findings have sparked renewed appreciation for the role of mitochondria in many intracellular processes coupled to synaptic plasticity and cellular resilience. New insights in depression pathophysiology are revolving around the impairment of neuroplasticity. Mitochondria have potential role in ATP production, intracellular Ca\(^2+\) signalling to establish membrane stability, reactive oxygen species (ROS) balance and to execute the complex processes of neurotransmission and plasticity. So understanding the various concepts of mitochondrial dysfunction in pathogenesis of depression indubitably helps to generate novel and more targeted therapeutic approaches for depression treatment.

Objective: The review was aimed to give a comprehensive insight on role of mitochondrial dysfunction in depression.

Result: Targeting mitochondrial dysfunction and enhancing the mitochondrial functions might act as potential target for the treatment of depression.

Conclusion: Literature cited in this review highly supports the role of mitochondrial dysfunction in depression. As impairment in the mitochondrial functions lead to the generation of various insults that exaggerate the pathogenesis of depression. So, it is useful to study mitochondrial dysfunction in relation to mood disorders, synaptic plasticity, neurogenesis and enhancing the functions of mitochondria might show promiscuous effects in the treatment of depressed patients.

Keywords: Depression, Mitochondrial dysfunction, Reactive oxygen species, neurotransmitter, Mitochondria, electron transport chain.

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INTRODUCTION

Mitochondria have pivotal role in energy production via metabolism of lipids, steroids and proteins, and in maintenance of cellular stability via modulation of Ca\(^2+\) levels, maintaining the levels of ROS and regulation of apoptosis [1]. Thus mitochondrial dysfunction not only hampers cells to meet energy requirement but might also be involved in the impairment of neuronal communication and cellular resilience, which prop up mood disorders and psychotic disorders [2, 3]. New hypothesis of mood disorder is revolving around the concept of neuroplasticity mainly depression and bipolar disorder. Neuroplasticity refers to the brain plasticity or brain malleability encompasses both synaptic plasticity and non-synaptic plasticity. It is the process of adapting neurons in response to changing internal or external environment by causing changes in neural pathways and synapses. Synaptic plasticity involves synaptogenesis, growth of axons and dendrite and removal of unnecessary connections between neurons. In neuroplasticity, mitochondria play an important role and it is well studied that stress leads to structural and functional impairment in various brain regions of depressed patients that lead to impaired neuroplasticity [4]. Neurogenesis i.e. the formation of new neurons is believed to be an important mechanism of brain plasticity under physiological conditions and in brain repair after injury [5, 6]. In various studies it has been consistently observed that different types of stresses decrease the adult hippocampal neurogenesis and lead to depression. During neuronal development, mitochondrial biogenesis occurs at higher rate as increased mitochondrial genome and mitochondrial protein is required for neuronal differentiation [7-9]. In this context, it is interesting to note that mitochondrial dysfunction might have an important role in impaired adult hippocampal neurogenesis in depression.
Mitochondrial Dysfunction in Depression

Mitochondria are the only organelle in the cell that has its own DNA called mtDNA (mitochondrial DNA). mtDNA codes only 13 protein subunits of electron transport chain (ETC) and rest mitochondrial proteins are coded by cellular DNA genes. It has well known function of producing cellular energy through citric acid cycle and oxidative phosphorylation (OXPHOS). Mitochondria are the power house of the cell producing ATP (adenosine triphosphate) by efficiently utilising oxygen and substrates like glucose and pyruvate. It fulfills 92% energy demand of body via OXPHOS. It consists of two membranes, an outer membrane and an inner membrane. Inner membrane is folded to form cristae on which the machinery for synthesis of ATP i.e. subunits of ETC are assembled in line. ETC consists of five protein complexes which work coherently to produce ATP at complex V. The three ETC chain complexes i.e. I, III and IV transfer H⁺ to the intermembrane space in mitochondria to produce electrochemical gradient for the synthesis of ATP [25, 26]. Human brain consumes 25% of total body glucose requirement. ATP, the cellular energy fuel is essential for the normal processing of neuronal processes including maintenance of ion gradients across neuronal membrane, accumulation of neurotransmitters in the vesicles, release of neurotransmitter and trafficking of receptors and ion channels to and from the cell surface [27, 28]. Brain stores only small amount of glycogen so it needs constant supply of glucose. Stressed neurons are unable to upregulate glycolysis and on inhibition of mitochondrial respiratory chain, neurons die rapidly and astrocytes use glycolytically derived ATP [29].

Reactive Oxygen Species

Reactive Oxygen Species (ROS) have significant role in pathogenesis of number of diseases; they are also mediators of several physiological processes. Mitochondria are the main intracellular organelle producing ROS [30]. Each complex of mitochondrial ETC has its own function but works in association with others. During electron transfer process O₂⁻ is produced at complex I and III. Mitochondrial superoxide dismutase (SOD) converts O₂⁻ into H₂O₂ and O₂. ·OH plays an essential role in cell physiology by activating guanaylate cyclase and formation of second messenger, cGMP and activation of NF-kB by H₂O₂. H₂O₂ gets converted into water by glutathione peroxidase and catalase in cytosol [31]. O₂−, highly reactive species form highly reactive molecule on interaction with nitric oxide having high oxidant and nitrating property, peroxinitrite which impairs enzymatic function tyrosine residues and results in decreased production of monoaminergic neurotransmitters and other aminergic compounds [32]. H₂O₂ can also react with Cu²⁺ and Fe²⁺ ions, resulting in formation of highly reactive free radical OH− that cause oxidative damage to carbohydrates, lipids, proteins, mtDNA and mitochondrial membranes, resulting in functional deficits and death. Monoamine oxidase A and B, located on the outer mitochondrial membrane metabolise serotonin, dopamine and noradrenaline following formation of free radicals [33]. In spite of formation of ROS, mitochondria also produce several protective antioxidant molecules, such as creatinine, coenzyme Q10, niconitamide and glutathione protecting neurons from various deleterious effects of oxidative reactions. Not only this, studies have shown that ROS have modulating role in synaptic plasticity, learning and memory formation [34, 35].

Mitochondria as Power House

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Ca\textsuperscript{2+} regulation

Mitochondria play an important role in calcium homeostasis, a principal second messenger that is involved in the regulation of neurotransmission and short and long term neuroplasticity in the brain. Both mitochondria and endoplasmic reticulum (ER) are involved in sequestering of intracellular Ca\textsuperscript{2+}. Outer mitochondrial membrane is permeable to Ca\textsuperscript{2+} and inner mitochondrial membrane has Ca\textsuperscript{2+} uniporters that facilitate inward movement of Ca\textsuperscript{2+} and Na\textsuperscript{+}/Ca\textsuperscript{2+} and H\textsuperscript{+}/Ca\textsuperscript{2+} antiporters that facilitate outward movement of Ca\textsuperscript{2+} [36]. The basal intracellular Ca\textsuperscript{2+} levels is very low compared to the extracellular space and ER, and even multiple, small changes in the concentration level result in repetitive partial mitochondrial depolarisation. Mitochondria have several mechanism for regulation and restoration of changes in Ca\textsuperscript{2+} levels. Mitochondrial matrix Ca\textsuperscript{2+} concentration is increased equivalent to increase in cytosolic Ca\textsuperscript{2+} levels and in situation of high energy demand and gets lowered in low cytosolic Ca\textsuperscript{2+} levels and high ATP/ADP ratios. In addition to the sequestering of cytocytic Ca\textsuperscript{2+}, mitochondria also contribute to the time, rate and amount of neurotransmitter release from presynaptic sites. A high concentration of Ca\textsuperscript{2+} in the mitochondrial matrix stimulate the activity of dehydrogenases i.e. pyruvate dehydrogenase, isocitrater dehydrogenase and \(\alpha\)-ketoglutarate dehydrogenase of the krebs cycle and complex V resulting in increased production of ATP [37]. At extremely increased Ca\textsuperscript{2+} uptake into mitochondrion results in cessation of ATP formation whereas increased rate of activation of mitochondrial permeability transition pore (mPTP). The increase permeability of the mitochondrial membranes to molecules of molecular weight less than 1,500 Daltons leads to mitochondrial swelling, cytochrome c release, and apoptosis activation [38-39].

Apoptosis

There are three forms of cell death i.e. apoptosis, necrosis and autophagic cell death. Apoptosis is the programmed cell death occurs in the neurons during development and adult cell turnover [40]. Apoptosis is essential to prevent growth of cancerous cells but uncontrolled apoptosis leads to neurodegeneration in post-mitotic cells such as neurons. Apoptosis is complex and energy dependent process in which dying cell itself involved in it. Morphologically, it is characterised by cytoplasmic shrinkage, chromatin condensation and rapid phagocytosis by neighbouring cells [41]. In normal physiological conditions apoptosis removes those neurons and glia which are unable to make neuronal connections. New neurons are produced in large number during adult neurogenesis and these neurons compete for making connections with other neurons and those neurons which have neither electrical activity nor neurotrophic support die by apoptosis [42]. Two biochemical pathways are involved in apoptosis i.e. intrinsic and extrinsic pathway, mitochondria play an important part in both the apoptotic pathways. Both pathways are activated by different stimuli but ultimately result in activation of proteases and cell death [43]. In extrinsic pathway, engagement of death receptors such as cluster of differentiation 95 (CD95), TNF-\(\alpha\)I, on the cell surface recruits Fas-associated death domain protein (FADD) which further triggers caspase-8 activation. Activated caspase-8 upon release in cytosol cleaves and activates other caspases and cleaves BH3 interacting domain death agonist (BAD), which then translocates to the mitochondria and initiates the caspase cascade which in turn triggers the executioner caspases. This ultimately leads to outer mitochondrial membrane permeabilization, leakage of apoptogenic factors and ultimately apoptosis [44]. In intrinsic pathway, mitochondria have significant role and is predominant in neurons. High levels of Ca\textsuperscript{2+} or ROS, damage caused by irradiation and activation of pro-apoptotic B-cell lymphoma protein 2 (BCL-2) family proteins, oxidative stress, over activation of glutamate receptors are the examples of stimuli that trigger activation of various kinases that lead to the loss of mitochondrial membrane potential. This occurs due to release of intermembrane cytochrome c into the cytoplasm. Cytochrome c then binds to cytosolic protein, apoptotic protease-activating factor 1 (Apaf-1) which further recruits and activates caspase 9 and results in the formation of a multisubunit enzyme complex known as the apoptosome [45, 46]. Activated caspase 9 then activates other executioner caspases like caspase 3 and 7 [47]. Similarly to cytochrome c, two intermembrane proteins, Smac/ DIABLO and Omi/HtrA2 inactivate IAP (inhibitors of apoptosis proteins), preventing caspase -3 and -9 inhibition. Once mitochondrial membrane integrity is impaired i.e. on mitochondrial membrane permeabilization (MMP), apoptosis inducer factor (AIF) and endonuclease G move to the nucleus and mediate chromatinolysis. All these processes lead to apoptosis by disabling repair processes, terminating cell cycle progression, inactivating inhibitors of apoptosis, mediating cytoskeletal and nuclear disassembly and activates cell phagocytosis [48-50]. Indeed neuronal apoptosis is essential in the early stages of development and during adult neurogenesis as it leads to the removal of damaged, unresponsive networks and also facilitates the synaptic plasticity and long-term potentiation.

Mitochondria and Neurogenesis

Neurogenesis is the process of formation of new neurons in which neural stem cells proliferate and then differentiate into neurons. Clear evidences from studies have shown that neurogenesis is prominent in the dentate gyrus of hippocampal region [51-55]. In depression, it has been shown that there is decreased hippocampal neurogenesis which leads to the formation of faulty neuronal networks and hence impairs adaptation to the changed milieu. During neuronal development, mitochondrial biogenesis occur at higher rate which leads to increase mitochondria number per neuron. Vayssiere, Cordeau-Lossouarn, Larcher, Basseville, Gros and Croizat [56] showed that inhibitor of mitochondrial protein synthesis, chloramphenicol prevents neuronal differentiation whereas oligomycin, mitochondrial ATP synthase inhibitor have no effect on the differentiation process. This suggests that increased mitochondrial genome and mitochondrial protein is required for neuronal differentiation [7-9]. Mitochondria can freely translocate with in the neuron. During neurite outgrowth the velocity at which mitochondria move decrease as the neurite outgrowth slows down and synaptogenesis occurs [57]. In the process of axogenesis, number of mitochondria increase in the neurite destined to become axon and decrease in neurite destined to become...
Mitochondria and Synaptic Plasticity

The long term changes in strength and activity of neuronal connections in response to stress and other aversive stimuli is called synaptic plasticity that involves structural and functional plasticity. In neurons, mitochondria have major function in regulation of developmental and synaptic plasticity [58]. The mitochondria in presynaptic axon terminals and postsynaptic dendritic spines differ in behaviour and functional properties. More than twice the number of mitochondria in the axon is motile as compared to the mitochondria in dendrites of cultured hippocampal neurons. Mitochondria in dendrites are highly charged and more metabolically active than in axons [59]. Studies have shown, mitochondria play an active role in synaptic plasticity [60, 61]. During synaptogenesis, the structural plasticity of dendritic spines is dependent on the number of mitochondria present in dendrites. Impairment of dynamin-like GTPases Drp1, mediate mitochondrial fission and OPA1, mediate mitochondrial fusion or over expression of dominant-negative form of Drp1 (A38K) reduces the number of dendritic mitochondria and impairs synaptogenesis whereas on treating cells with creatine increases dendritic mitochondrial mass and activity and approximately double the number of synapses [62-65]. Thus mitochondria have an imperative role in structural synaptic plasticity. The function of presynaptic neuronal axons and postsynaptic neuronal dendrites are highly dependent on the changes in the intracellular Ca²⁺ levels as a result of metabolic and oxidative stress. Mitochondria act as Ca²⁺ sensor and take up Ca²⁺ after synaptic stimulation and thereby playing a role in neuronal transduction [66]. Mitochondria have also been involved in LTP, a form of synaptic plasticity. It has been shown in studies that ROS are essential for the induction of LTP [67, 68]. Various studies have shown that superoxide, hydrogen peroxide and nitric oxide are involved in hippocampal synaptic plasticity. These ROS increase the activity of protein kinase C, extracellular-regulated kinase 2, protein tyrosine kinases ryanodine receptor type 3, and decrease the activity of protein phosphatases 2A and 2B, and a protein tyrosine phosphatase [69-71]. Moreover, it has been found in in-vitro study that on treatment with scavengers of ROS attenuated LTP in hippocampus slices and in in-vivo study where hippocampus of transgenic mice overexpress Cu/Zn superoxide dismutase [72, 73]. Excessive ROS have detrimental effect on the functions of brain. Similarly, excessive Ca²⁺ levels lead to the activation of pro-apoptotic factors and activation of apoptotic process lead to LTD (long term depression), another form of synaptic plasticity. Thus, a balance between LTD and LTP in the brain is essential for mental health [74]. In addition to the changes in the movement of mitochondria within axons and dendrites, changes in other mitochondrial function such as Ca²⁺ regulation, energy metabolism and ROS production also play important role in synaptic plasticity. Emerging findings also suggest that mitochondria exhibit some the effects of BDNF and glutamate on synaptic plasticity. Markham, Cameron, Franklin and Spedding [75] showed increase in glucose utilization in cultured cortical neurons in response to enhanced energy demand and increase in the respiratory control index in rat brain mitochondria on treatment with BDNF [75, 76]. Regulation of various processes involved in synaptic plasticity by mitochondria enlightens its essential role in synaptic plasticity.

Mitochondrial Dysfunction in Depression

Brain is highly active organ with high energy consumption and organs with highest energy demand are more susceptible to the deleterious effect of reduced energy production. Mainly neurons obtain energy from mitochondrial OXPHOS. Neuron obtains energy through glycolysis but brain is unable to store glycogen so constant supply of glucose is needed. When oxidative phosphorylation is compromised, neurons are unable to meet their energy demand and alteration in their various physiological functions occurs due to reduced ATP production. Studies have shown that chronic mild stress inhibits mitochondrial OXPHOS, dissipates mitochondrial membrane potential and damages mitochondrial ultrastructure in various brain regions including hippocampus, cortex and hypothalamus of mice [77, 78]. These observations lead to the conclusion that most probably depression is caused by energy impairment in the brain due to mitochondrial dysfunction [79, 80]. In support of this notion, studies have shown that patient suffering from depression showed reduced glucose utilization in the PFC, anterior cingulated gyrus and caudate nucleus [81-83].

The relationship between social–psychological and physical stress and the development of depression in susceptible individuals has long been established [84]. Rodents subjected to chronic mild stress show depression like symptoms [85]. Major depression is associated with reduced hippocampal volume. Both structural and functional changes within several brain regions, including the hippocampus, amygdala and prefrontal cortex, resulting in faulty connections between anterior cingulate cortex (ACC), PFC, hippocampal, and amygdaloid regions, ends up with constant alterations in neuroplasticity [86, 87].

Hyperactivity of HPA axis (Hypothalamic–pituitary–adrenal axis) in depression is associated with increased glucocorticoid (GC) levels both in CNS and peripherally [88, 89]. GCs play biphasic role in regulating mitochondrial functions. Cultured neurons on acute exposure to either low or high levels of glucocorticoids are markedly increase mitochondrial activity. GCs bound to the glucocorticoid receptors can translocate to mitochondria and inhibit the formation of Bax- containing pores on the mitochondrial outer membrane by forming a complex with the anti-apoptotic protein Bcl-2, and reduce the release of calcium and cytochrome C from the mitochondria and ultimately inhibit apoptosis [31]. Whereas chronic elevation in GCs level cause kainic acid induced toxicity of cortical neurons. Prolonged exposure to GCs also cause respiratory chain dysfunction, increased ROS generation, mitochondrial structural abnormalities, apoptosis and cell death in skeletal muscle cells and hippocampal neurons [90, 91].
Stress also increases the levels of proinflammatory cytokines including IL-1β, IL-6, and TNFα and there are ample evidences showing that debilitating major depression is accompanied by activation of immune, inflammatory, oxidative, increased HPA-axis and nitrosative stress pathways. Animals subjected to different stresses like psychological stress, immobilization stress, inescapable shock, physical restraint, open field stress or conditioned, aversive stimuli showed increased pro-inflammatory cytokines and nitric oxide in the brain and plasma [92-94]. TNF-α, a pro-inflammatory cytokine, suppresses mitochondrial complexes I and IV and pyruvate dehydrogenase activities and induces mitochondrial damage [95]. IL-6 stimulates increased ROS production in the brain [96] and nitric oxide is the potent inhibitor of mitochondrial cytochrome c oxidase [97]. Cytokines also induce activation of pro-apoptotic members of the BCL-2 family and activation of the caspase cascade. It has been shown that unpredictable stress reduces the expression of Bcl-2 mRNA in the rat limbic structures, an anti-apoptotic endogenous membrane protein that promotes neuritogenesis and axon regeneration [98-100]. BAG-1 (Bcl-2 associated athanogene) is a gene that binds to Bcl-2 and enhances the anti-apoptotic effects of BCL-2. Chronic mild stress has been shown to reduce the expression of BAG-1 [101]. This ultimately leads to the activation of caspases and BCL-2-associated X protein (BAX) and BCL-2 antagonist/killer (BAK) in the mitochondria which further alters membrane permeability, ends up with neuronal death. The production of ATP and Ca²⁺ buffering is also an important function of presynaptic mitochondria in order to maintain neuron-neuron communication by controlling synaptic vesicles mobilization and recycling process. Vesicle cycling and neurotransmission is highly dependent on mitochondrial function as many steps involved in it are ATP consuming. Mobilization of synaptic vesicles to the active sites of synapse and endo/exo cycling pool process is dependent on ATP. Synaptic vesicles propulsion is dependent on the activation of myosin ATPase complex which results in the translocation of actin filaments. Upon activation ATP gets converted into cAMP by phosphodiesterase and adenylatecyclase. cAMP then activates PKA kinase, resulting in the mobilization of synaptic vesicles to the active site [102]. Similarly in a mammalian study, it has been observed that impairment of interaction of syntabulin with

**Fig. (1).** Exposure to different stressful situations leads to increase ROS production, increased levels proinflammatory cytokines, increased nitrosative stress and decrease in antioxidant enzymes level which ultimately leads to reduce OXPHOS, activation of apoptotic pathway and causes mtDNA damage as consequence of which there is decreased mitochondrial biogenesis, increased ROS production, apoptosis of neuronal cells, impaired translation of mitochondrial ETC complex proteins and decreased ATP production which results into mitochondrial dysfunction. Neurogenesis, synaptic plasticity and neuronal transmission, the important parameters for successful adaptation to the stressful conditions are also compromised due to mitochondrial dysfunction hence also play an important role in major depression.
kinesin impairs neurotransmitter release due to inhibition of mitochondria translocation at the active site of synapse and is reversed by exogenous ATP addition [103]. Thus, defects in ATP production and disturbed mitochondrial translocation to the active site of synapse lead to the local ATP depletion and defective neuronal transmission [104, 105].

An imbalance between ROS production and antioxidant defence mechanism may represent the one factor influencing the etiology of depression. It is reported that chronic unpredictable mild stress impairs the endogenous antioxidant status in the brain. Various studies showed that chronic mild stress induced depression in rodents lowers the concentration of brain glutathione, total antioxidant capacity, superoxide dismutase and catalase [106-108]. Hence, there is increased oxidative stress in depression. As mitochondria are the main site of ROS production in the cell, they are highly susceptible to oxidative damage. Lipids, proteins, oxidative phosphorylation enzymes and mtDNA in mitochondria are more susceptible to oxidative damage [109]. Thus, mitochondrial functions are impaired and aggravate oxidative stress, increase in intracellular Ca²⁺ levels, damage or deletions of mtDNA, alterations in fusion/fission and morphology of mitochondria and finally to neuronal death [110]. Further mitochondrial dysfunction is compromised by increasing energy demand for cellular repair process and in this way mitochondrial damage causes additional damage [111].

CONCLUSION

The exact pathogenic mechanisms behind depression are still not perfectly known but increasing evidences suggest that mitochondrial dysfunction may be involved in various psychiatric diseases. Regulatory role of mitochondria in synaptic strength and cellular resilience in neuronal circuits mediate complex, high-order brain functions and implicate mitochondrial dysfunction in the pathogenesis of mood disorders. Thus, enhancing mitochondrial functions appears to be reasonable for increasing mitochondrial function and preventing or alleviating the burden of many stress-related disorders. Targeting mechanisms of mitochondrial dysfunction due to its special impact on energy metabolism, synaptic plasticity and neuronal survival, may be an important avenue for development of new mood-stabilizing agents. The mitochondrially-targeted therapeutics that have reached clinical trials so far have produced encouraging but largely inconclusive results. Uridine showed statistically significant improvement in depressive symptoms in Phase IIa trial whereas Phase IIb study showed negative results [112]. Coenzyme Q10, acetyl L-carnitine and α-lipoic acid are also being studied for treatment of Bipolar disorder (ClinicalTrials.gov identifiers NCT01390389 and NCT00719706). Increasing understanding of mitochondrial dysfunction generated exciting preclinical data that show association of mitochondrial dysfunction with neurodegenerative disorders but less clinical evidences are available so further work is needed to shed light on the role of mitochondrial dysfunction in the pathogenesis of neuronal disorders.

CONFLICT OF INTEREST

The authors confirm that this article content has no conflict of interest.

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