Chemism and the Role of Endocannabinoids in Physiological Processes

ABSTRACT

Despite being known for 5000 years, after the records of imperial Chinese doctors, cannabinoids as a subject of scientific research experienced its rise after 1964, when delta nine tetrahydrocannabinol (Δ 9-THC) by Israeli scientists was identified. This was followed by the discovery of endogenous ligand / endocannabinoids, as well as receptors CB1 and CB2.

In a broader sense, endocannabinoids act as neuromodulators and immunomodulators. They are included in the various physiological processes such as: the occurrence of pain, cognition, memory formation and neuroplasticity, physical activity, respiratory processes, appetite regulation, control and heart rate, nausea and emesis, intraocular pressure, inflammatory and immune processes (antigen recognition).

Key Words: Endocannabinoids, anandamide, cannabinoid receptors.

Introduction

Endocannabinoids were named after a plant cannabis or fitocannabinoids. Studying fitocannabinoids has resulted in the development of the most important physiological endocannabinoid system (ECS) that is important for the establishment and maintaining homeostasis in humans.

In 1964, dr. Raphael Mechoulam, dr. Lumir Ondrej Hanus and their followers at the Hebrew University in Jerusalem identified Δ (delta) -9-tetrahydrocannabinol (THC) in cannabis. The following research led to the development of a receptor on which THC is connected as well as the development of endocannabinoid and endocannabinoid system.

Endocannabinoids are endogenous agonist/ligand of cannabinoid receptors CB1 and CB2, which are produced in mammalian tissues.

The first endogenous cannabinoid anandamide (arachidonylthanolamide or AEA) was isolated and described by Czech chemist Ondrej Lumir Hanus and American molecular pharmacologist William Anthony Dekane in 1992. The name itself was given according to the Sanskrit word ananda which means blissfulness. Shortly after the discovery of the first one, the other endogenous cannabinoid, 2-arachidonoylglycerol (2-AG), was developed by Shimona Ben-Shabat, one of the first students of the above mentioned doctors.

Picture 1. Anandamide (AEA) (3).
Endocannabinoids are derived from arachidonic acid and they can be amid, ester or eter with long unsaturated fat acids. According to their own chemism, they are hydrophobic molecules and hence they do not migrate through the body so their engagement is local, surrounded by the space that is close to synthesis molecules endocannabinoids.7,8

The most examined endocannabinoids (Picture 1 and 2) are arachidonyletanolanid (anandamide, AEA) and 2- arachidonyl-glycerol (2-AG). The theory, according to which their synthesis appears when necessary, was refuted and it was proved that anandamide was and is contained in some cells. The other representatives of endocannabinoid are: 2-arachidonyl-glycerol (noladin), O- arachidonyletanolanim (virodhamine), N-arachidonylidopamine and others.

Synthesis of anandamide is carried out paralelly in a few different ways upon which are contained different enzymes (phospholipase D, phospholipase C, α, β, hydrolase 4 and different phospholipases). Precursor for the synthesis of anandamide is a membrane phospholipid N- arachidonyle phoshatidyl- ethanolamine.6

Synthesis of anandamide in neuron is stimulated by a binding of a neurotransmitter that is released from presynaptic neuron in an adequate inotrope or metabotropic receptor on postsynaptic neuron.

This process causes increased cytosolic free calcium-ions concentration in postsynaptic neuron which represents a stimulant for synthesis and releasing endocannabinoids from their precursors in a membrane. The released endocannabinoids are bound to receptors CB1 on presynaptic membrane. After that, activated receptors inhibitate potassium canals that are dependent on power.

Due to the increased diffusion of potassium ions, depolarization of presynaptic membrane is decreased. The consequence is the inhibition of neurotransmitters release such as glutamate, dopamine and γ- aminobutyric acid (GABA).7,8

Termination of biological results of anandamide is carried out in two parts. In the first part, anandamide migrates into the cell, while in the second part, it is decomposed with the help of hydrolase amid fatty acid FAAH (Fatty Acide Amide-Hydrolase).9 Bearing in mind that anandamide is a lipofin molecule, it can migrate into the cell via passive diffusion in the direction of concentrated gradient. The transport anandamide in the cell is also enabled via selective transport molecule called AMT (anandamide membrane transporter), which is placed in the membrane plasma and acts reversible. There are two other known mechanisms by which it is possible to endorse anandamide in the cell, that is, with intracellular membrane vesicul and with the help of endocytosis.10-14

Using transgenic animal models, without genes for receptors CB1 and CB2, other receptors on which endocannabinoid is carried out were gradually developed. Receptors TRVP, (transient receptor potential cation channel subfamily V member vaniloid receptor) participate in the regulation of body temperature and system of nocicepsys that medites signals to pain stimuli. Receptors GPR55 (protei-coupled receptor) has 13% of homology with receptors CB1 and 14 % of homology with receptor CB2. It is placed in the brain, liver, spleen, intestines, fetal tissues and placenta. It is a very serious candidate to be named receptor CB1.15,16

The role of endocannabinoid in physiological processes

In wide range, endocannabinoids work as a neuromodulator and immunomodulator. They are included in different physiological processes such as: pain, cognitive processes, memory formation and neuroplasticity, motoric activities, endocrine processes, regulation appetite, control and puls, nausea, intraocular pressure, inflammatory and immune processes (antigen).17,18

Endocannabinoids are intracellular transmitters (vectors) of signals close to the neurons in synopsis. Since they are lipophilic molecules, they are not contained in intracellular vasculun but after synthesis, they become a part of cellular membrane. It is characteristic for them to participate in retrograde signalling between neurons, which means that the signal, instead from presynaptic neuron, travels to postsynaptic neuron in another way (Picture 3). Endocannabinoids are released in the synaptic cut from the postsynaptic neuron and act on presynaptic nerve endings. Activation of cannabinoid receptors
on presynaptic neuron for the small amount of time inhibates the release of the second neurotransmitter. The final result is dependent on the type of neuron which is inhibited from cannabinoid such as, for example, inhibition of excitatory neurotransmitters such as glutamate leads to inhibition of an excitatory neuron, while inhibition of releasing inhibitory neurotransmitter GABA (gamma-aminobutyric acid) leads to the increasement of excitability.\textsuperscript{19}

**Picture 3. Schematic diagram of endocannabinoid activity in presinapse and retrogressive modulation of releasing neurotransmitters glutamate and GABA (gamma-aminobutyric acid), AEA (arachidonylethanolamide), 2-AG (2-arachidonyl-glycerol) (20).**

**Endogenous cannabinoid receptors**

The first proof of the presence of cannabinoid receptors was the evidence that THC inhibates adenyl cyclase, after which studies that used radioligands followed.\textsuperscript{18,20} The first cannabinoid receptor CB\textsubscript{1} was cloned in 1990 from the cell of a mouse brain, while the other cannabinoid receptor CB\textsubscript{2} was cloned in 1993 from human cells of promyelocytic leukemia HL-60.\textsuperscript{21,22}

Cannabinoid receptors are divided in two types - CB\textsubscript{1} and CB\textsubscript{2}. They are rated as high affinity 7-transmembrane receptors that are connected with G protein. Activation of receptors causes the induction of different intracellular processes: inhibition of adenyl cyclase and, as a consequence, concentration of CAMP decreases (adenoid cyclic 5- monophosphate), activation with mitogen- activated protein kinase ERK (extracellular signal-regulated kinase), amino terminal kinase, activation of phosphatidylinositol 3-kinase, degradation of sphingomyelin, as well as the ceramide occurrence.\textsuperscript{23,24}

Receptors CB\textsubscript{1} and CB\textsubscript{2} have just 44% conguence with the chain amino acids while in the dome of binding, they have more similarity in sequences of amino acids (68%), which is the reason why some ligands do not make difference in binding to CB\textsubscript{1} or CB\textsubscript{2} receptors. It is interesting that human receptors CB\textsubscript{2} have high similarity to other animal types ( monkeys 100%, rats 97%, mice 96%).\textsuperscript{25}

**Receptors CB\textsubscript{1}**

Receptors CB\textsubscript{1} are present in the central nervous system as well as in the periphery tissues. In the brain, they are in the parts that control motor activities (basal ganglia and cerebellum), memory and cognitive functions (cortex and hippocampus), emotions (amigdala), sensory perceptions (thalamus), autonomy and endocrine functions (hypothalamus, pons and medul). Distribution of CB\textsubscript{1} receptors in the brain is in accordance with famous pharmacodynamic results regarding the effects of cannabinoid on memory, cognitive activities, pain and movement. Low concentration of receptors CB\textsubscript{1} is present in a brainstem, extended spinal cord and hilaminus. The above mentioned explains why cannabinoids do not cause mortality due to dangerous acts on vital physiological functions in cases of ingestion of a very high dose of cannabinoid. According to the data published in American Scientist in December 2015, mortality associated with the use of cannabis in the USA was around 0, as well as in 2014 and, in contrast to alcohol, which caused 30700 deaths in 2015.\textsuperscript{27} On the periphery, they are located in testicles, vascular endothelium, spleen and periphery nerves.\textsuperscript{1,12,23,28}

**Receptors CB\textsubscript{2}**

In comparison with CB\textsubscript{1} receptors, CB\textsubscript{2} receptors are primarily manifested in the cells of the immune system and they participate in modulation of functional aspect of the immune system. The large number of receptors CB\textsubscript{2} is expressed in lymphocytes B and in natural cells killers (natural killers NK). Additionally, they are present in tonsils, spleen and in lymph nodes. Some cannabinoids act on validus receptors and T-type calcium channels. It is generally considered that psychoactive cannabinoid results are dispatched through CB\textsubscript{2} receptors, while immunomodular ones are dispatched through CB\textsubscript{2} receptors.\textsuperscript{21,28,30}

**Conclusion**

The oldest records regarding the medical use of cannabis are known since the time of Chinese imperial herbalist Shen Nung 5000 years ago, who recommended cannabis for curing malaria, beriberi, constipation, rheumatological pains, shortage of concentration and woman problems. The famous doctor of that time in China, Hua Tuo, used the mixture of cannabis resin and
wine as analgesic in cases of surgery. Medical use of cannabis is also well known in all old civilizations such as India, Mesopotamia and Ancient Egypt.\textsuperscript{30,31} Although cannabinoids have been used for centuries, the range of studies that examines the consequences of cannabinoid has developed in the last 50 years, after the development of cannabinoid receptors and their endogenous ligand. The research is not only conducted in the field of basic mechanism of cannabinoid activity but in the developing pharmacological and therapeutic results of cannabinoid. Until now, there are more than 150000 studies connected to the theme of endocannabinoid systems, endocannabinoid as well as pharmacological and therapeutic activity of phyto-cannabinoid. The fact that more and more countries all over the world legalize the use of medical cannabis speaks in favor of justification of clinical studies carried out with the usage of cannabinoid, as well as the validity of positive results that are obtained by these studies.

References

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Hemizam i uloga endokanabinoida u fiziološkim procesima

SAŽETAK

Iako poznati već 5000 godina po zapisima carskih kineskih ljeekara, kanabinoidi kao predmet naučnih istraživanja doživljavaju svoj uspon nakon 1964. godine kada je od strane izraelskih naučnika identifikovan delta девет tetrahidrokannabinol (Δ 9 THC). Nakon toga uslijedio je olakšaj endogenih liganda kao i receptora CB1 i CB2, na koji se vežu endokanabinoidi.

Endokanabinoidi u širem smislu djeluju kao neuromodulatori i imunomodulatori. Uključeni su u različite fiziološke proceze kao što su pojava bolesti, kognitivni procesi, formiranje pamćenja i neuroplastičnost, motoričke aktivnosti, endokrini procesi, regulacija apetita, kontrola i broj otkucaja srca, mučnin i povraćanja, intraokularni pritisak, inflamatorni i imunološki procesi (prepoznavanje antigena).

Ključne riječi: Endokanabinoidi, anandamid, kanabinoidni receptori.