Second messengers in endocrinology: a mini-review of the cyclic nucleotides

Abstract

In this mini-review the developments in the cyclic nucleotide “second messenger” system are discussed. Since the discovery in 1957 knowledge about the downstream effects of the cyclic nucleotides has increased enormously. The EPAC system and the newly discovered cyclic-di-monophosphate system will have promising therapeutic applications in the future and are new challenges.

Keywords: Second messenger, Cyclic nucleotides, EPAC system, Cyclic-di-monophosphate, cAMP, Cyclic nucleotide phosphodiesterases

Introduction

Second messengers are molecules inside cells that act to transmit signals from a receptor to a target. The term second messenger was coined upon the discovery of these substances in order to distinguish them from hormones and other molecules that function outside the cell as “first messengers” in the transmission of biological information.

Many second messengers are small and therefore diffuse rapidly through the cytoplasm enabling information to move quickly throughout the cell. As elements of signalling pathways second messenger scan serve to integrate information when multiple independent upstream inputs influence the rates of synthesis and degradation of the second messenger. In addition second messengers can have multiple downstream targets thereby expanding the scope of signal transmission.

A large number of second messengers have been characterized including cyclic nucleotides, hydrophobic molecules eg cyclic adenosine monophosphate or cAMP and cyclic guanosine monophosphate or cGMP (hydrophobic molecules), ions (eg Ca^{2+}), phospholipid derived molecules (eg., inositol triphosphate, phosphatidyl inositol ) and even gasses as nitric oxide ( NO), H_{2}S and carbon monoxide (CO).

The calcium ion Ca^{2+}, which can diffuse both through cytosol and across cellular membranes, has a critical role in the rapid response of neurons and muscle cells. At rest cells maintain a low concentration of Ca^{2+}, in the cytoplasm expending energy to pump these ions out of the cell. When activated neurons and muscle cells increase their cytoplasmic Ca^{2+}, concentrations by opening channels in the cell membrane which allow Ca^{2+}, ions outside the cell to enter rapidly. For the purpose of this mini-review only the cyclic nucleotides are discussed.

The cyclic nucleotides

The cyclic nucleotide cAMP is synthesized by adenylate cyclase enzymes which are downstream of G–proteins (guanine nucleotide binding proteins) and receptors. Cyclic AMP generated by adenyl cyclase has three major targets in most cells.

1. The cyclic AMP–dependent protein kinase (PKA)
2. cAMP regulated guanine nucleotide exchange factors termed EPACs (exchange protein directly activated by cAMP )
3. Via PKA phosphorylation by a cAMP response binding protein called CREB.

The most common downstream effector of CAMP is protein kinase A (PKA). PKA is normally inactive as a tetrameric holoenzyme (two catalytic and two regulatory units). The regulatory unit always blocks the catalytic center of the catalytic unit. The regulatory unit dissociates from the catalytic subunit. The free catalytic subunit interacts with proteins to phosphorylate Ser or Thr residues, thus providing a cellular response.

PKA can phosphorylate a diverse array of physiological targets such as metabolic enzymes and transport proteins and numerous regulatory proteins including other protein kinases, regulation of glycogen, sugar and lipid metabolism, ion channels and transcription factors.

Cellular transcription factors as the neutrophin BDNF (Brain Derived Neurotrophic Factor), tyrosine hydroxylase and many neuropeptides bind to certain DNA sequences called cAMP response elements (CRE) thereby increasing or decreasing the transcription of the downstream genes.

Protein synthesis by PKA directly activates CREB which bind the cAMP response element (CRE). The activated CREB protein then binds to a CRE region and is then bound to CBP (CREB binding protein) which co activates it allowing it to switch within genes on and off and altering the transcription.

CREB proteins have many functions in many different organs. CREB proteins in neurons are thought to be involved in short- and
long–term memory. CREB proteins also have an inhibiting effect on T–cell proliferation and its' effector effects via cAMP.

To integrate extracellular signals from membrane receptors with diverse intracellular signalling pathways the system uses cAMP–regulated guanine nucleotide exchange factors (GEFs). For that purpose small GTPases operate as binary switches that exist in GTP or in GDP ligand formation. They are involved in processes like cell adhesion, phagocytosis, apoptosis and gene expression.

The two GEFs regulated by cAMP are able to activate members of the Ras small GTPase family Rap1 and Rap2.

(EPAC1 and EPAC2) New paragraph? The EPAC pathway is involved in immune, lung and neuronal function. EPAC1 regulates the differentiation of monocytes to macrophages and subsequently controls cellular phagocytosis as well as the production of inflammatory mediators and apoptosis in human leukocytes. EPAC1 modulates integrin function most probably by barrier function and leukocyte adhesion and transmigration.

Mature lymphocytes with their antigen–specific receptors are in a permanent circulation from the blood stream to the lymphoid organs and represent the main mediators of the adaptive immune response. In B–cells signalling by the antigen–specific B–cell antigen– receptor complex (BCR) is a central step to maintain homeostasis and proper immune function with cAMP in the role of a central mediator. Expression of EPAC1 has been reported in B cell chronic lymphocytic leukemia.

EPACs represent a novel key effector in the lung due to their capacity to modulate airway inflammation and proliferation and as part of the signalling cascade triggered by beta 2–adrenoceptor agonists that target cAMP to alleviate the symptoms of asthma.

It has been postulated that the developmental regulation of EPAC expression promotes axon growth and regeneration in the nervous system. Epac2 signalling inhibition induced early after–depolarization in ventricular myocytes in the heart. The underlying mechanism involves an increase in mitochondrial reactive oxygen species (ROS) and activation of the late sodium current (i NaLate)? Inhibition of EPAC2 caused ventricular tachycardia, torsades de pointes and sudden death.

In the pancreas the production of the incretin hormone glucagon–like–peptide–1 (GLP–1) is stimulated by activation of PKA and EPAC2. In the brain EPAC2 may regulate synaptic plasticity and thus control higher brain functions such as memory and learning.

Pharmacological manipulation of cAMP levels exerts beneficial effects through the regulation of EPAC. For example EPAC2 selective agonists could promote insulin secretion from pancreatic beta cells whereas EPAC1 selective agonists may be useful in the treatment of vascular inflammation. In contrast EPAC1 and EPAC2 antagonists could both be useful in the treatment of heart failure.

Cyclic nucleotide phosphodiesterases form another family of important signalling proteins. Their activities are regulated via the rate of gene transcription as well as by second messengers (cyclic nucleotides or Ca2+) and interactions with other signalling proteins such as beta–arrestin and protein kinases. PDEs hydrolyze the cyclic 3,5–phosphodiester bound in cAMP and cGMP thereby terminating their action, Phosphodiesterase 4 (PDE4) is the predominant degrading enzyme expressed in inflammatory cells. Selective PDE4 inhibitors can play a therapeutic role in various inflammatory diseases as asthma, psoriasis, inflammatory bowel disease etc.

Cyclic AMP promotes the release of the anti–inflammatory mediators (eg IL–10) by immune cells. A decrease in PDE 4 increases a cAMP level which leads to increased transcription of genes that have CRE sites, including the gene for IL–10. PDEs are drug targets for the treatment of diseases as asthma, COPD, cardiovascular diseases as heart failure, atherosclerotic coronary and peripheral arterial disease and neurological disorders. Recently cyclic di–adenosine monophosphate was discovered (c–di–AMP). It is a broadly conserved messenger for bacterial growth and infection. However the molecular mechanisms of c–di–AMP signalling are still poorly understood.

Cyclic GMP

The signalling pathways that regulate the synthesis of cyclic GMP in cells include hormonal regulation of transmembrane guanylate cyclases such as via the natriuretic peptide receptor (ANP) and the activation of soluble guanylate cyclases by nitric oxide (NO). Unlike cAMP the second messenger cGMP has established signalling roles in only a few cell types. The synthesis is regulated by NO or by the natriuretic peptide hormones ANP, BNP (brain natriuretic peptide) and CNP (C–type natriuretic peptide). NO is synthetized from arginine and oxygen by the NO synthetase. It diffuses freely through the plasma membrane of the cells. It activates soluble guanyl cyclases to produce cGMP. NO synthesis by endothelial cells is stimulated by the neurotransmitter acetylcholine. The function of NO is blood vessel dilatation and smooth muscle relaxation.

Nitric Oxide Synthetase (NOS) exists in 4 isoforms: the neuronal type 1 isoform (nNOS), the inducible type 2 isoform (iNOS), the endothelial type 3 isoform (eNOS) and the mitochondrial isoform (mNOS). NOS is expressed in endothelium, cardiac myocytes, renal mesangial cells, osteoblasts and platelets and is involved with vasodilation (30) nNOS serves nerves and skeletal muscles and performs a role in cell communication and plays also a role in regulating blood pressure in healthy humans. iNOS is expressed in macrophages, Kupffer cells, neutrophils, fibroblasts and vascular smooth muscle cells and endothelium. It is involved in the immune defense against pathogens. mNOS regulates ATP synthesis in the mitochondrial matrix.

The transmembrane receptors for ANP, BNP and CNP are ligand–activated guanyl cyclases. Phosphorylation of serine residues in the kinase domain is important for activity. Dephosphorylation of these residues leads to desensitization of the receptor. After phosphorylation of the serine residues guanyl cyclase production is stimulated. Downstream reactions of cGMP are the activation of protein kinase G (PKG), gate keeping the ion channels and the formation of a cGMP modulated phosphodiesterase. Pharmacological important effects of elevated cGMP levels include modulation of platelet activation and relaxation of smooth muscle. PDEs hydrolyze the cyclic 3,5–phosphodiester bond in cAMP and cGMP thereby terminating their action.

While the natriuretic peptides use the guanyl cyclase system hormones like glucagon, epinephrine and parathyroid hormone (PTH) use the adenylate cyclase system. States of excess or low PTH production can be studied by measuring nephrogenous cAMP excretion and phosphaturia (TmPO4/GFR). States of end–organ resistance to PTH can be studied also in this way.

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The peptide hormones use the hormone receptor complex. Pharmacologic drugs use the ligand–receptor complex which is very similar to the hormone–receptor complex. However in this case the drug is the ligand.

**Selected drugs that stimulate cAMP formation are**

a. Salmeterol, a beta2 agonist used in asthma and COPD.

b. Haloperidol, a D2, D3, D4 antagonist used in schizophrenia.
   - Metoclopramide, a D2, 5 HT4 antagonists used in nausea and vomiting.

ii. Desmopressin, a V2 receptor agonist used in diabetes insipid us.

c. Drugs that decrease cAMP formation are for example:

   d. Metoprolol, a beta 1 antagonist used in angina pectoris, hypertension and heart failure.

c. Morphine, an u agonist? Used in pain relief.

f. Ibuprofen, a non–selective COX–inhibitor used in inflammation or pain.

g. Carbagoline, a D2 agonist used in Parkinson’s disease.

h. Drugs that use the NO–cGMP system are for example:
   - Nitroglycerin, isosorbide mononitrate. They are NO donors used in angina pectoris
   - Sodium nitroprusside used in hypertensive emergencies is a NO donor.

k. Niorandil, a K+ channel opener and NO donor used in angina pectoris.

l. Nifedipine, a NO releaser used in angina pectoris and hypertension.

**Synthetic natriuretic peptides are**

i. Nesiritide, a synthetic BNP that promotes vasodilatation, diuresis, natriuresis and used in heart failure.

ii. Ecadotril‒Neural Endopeptidase (NEP), catalyses BNP degradation, used in CHF.

**PDE inhibitors**

a. They act by modulating the levels of second messengers.

b. PDE 3 inhibitors, Amrinone, Milrinone used in CHF as vasodilators.

c. PDE 5 inhibitors, Sildenafil, Tadalafil used in erectile dysfunction. Produce vasodilatation by NO dependent elevation of cGMP in vascular smooth muscle.

d. Theophylline, Aminophylline, Non–selective PDE inhibitors used in asthma and COPD.

e. Pentoxifylline, a PDE 3 inhibitor with rheologic modifying properties. Improves the microcirculation.

f. Cilostazol, PDE 3 inhibitor that promotes vasodilatation and inhibits platelet aggregation.

g. Dipyridamole, a PDE 5 inhibitor. Inhibits thrombocytes aggregation. Prevents uptake and degradation of adenosine.

h. ANagrelide, a PDE 3 inhibitor. Used in essential thrombocythemia and Polycythemia Vera.

i. Anti–Spasmmodic as Dratuverine, inhibits PDE 4. Smooth muscle relaxant Used in irritable bowel syndrome, renal and biliary colics, dysmenorrhea and acceleration of labor.

**Conclusion**

Since the discovery of cAMP in 1957 by the late 1971 Noble Prize winner Sutherland the knowledge about the downstream effects of the cyclic nucleotide “second messenger” system has increased enormously. In particular the EPAC system has the potential for many yet unknown therapeutic applications. The discovery of the new cyclic–di–adenosine monophosphate system is another new promising challenge. Cell biology has done a great job in cyclic nucleotide research and clinical pharmacology will benefit without doubt.

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**Conflicts of interest**

The author declares that there are no conflicts of interest.

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**References**


