Tendencies and perspectives of central alpha2-adrenomimetic application in medico-biological research

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ABSTRACT

The pharmacodynamic effects of the administration of a2-adrenergic agonists both in a monovariant and in combination with drugs of other pharmacological groups are considered. Based on analysis of safety nonclinical studies the characteristics of main physiological effects of a2-adrenergic receptors as well as physiological effects of a2-agonists on various organs and systems are presented. For the determination of tendencies and directions in research of central a2-AM (dexmedetomidine) the analysis of bibliographical data, accumulated and extracted from Medline database with 5 year time-filter (VOSviewer, 1.6.11 version) has been carried out. For the further research of central a2-adrenomimetics and their application in clinical practice the following perspective directions have been determined: the study of effects and mechanisms of cytoprotective and antioxidant action, the study of the use of drugs in a monovariant and in combinations for the development of analgesic drugs, anesthesia and development of combined formulations with a delayed release of antagonists designed to mitigate side effects.

KEYWORDS

Alpha2-adrenomimetics, a2-agonists, dexmedetomidine, combinations, antioxidant, cytoprotection, lipid peroxidation.

Тенденции и перспективы применения центральных альфа2-адреномиметиков в медико-биологических исследованиях (обзор литературы)

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РЕЗЮМЕ

Рассмотрены фармакодинамические эффекты при введении a2-адреномиметиков как в моноварианте, так и в комбинации с препаратами других фармакологических групп. На основе анализа доклинических исследований безопасности представлена характеристика основных физиологических эффектов a2-адренорецепторов, а также физиологических эффектов при воздействии a2-агонистов на различные органы и системы. Для определения тенденций и направлений исследования центральных a2-АМ (по дексмедетомидину) выполнен анализ библиографических данных, отобранных в базе MEDLINE с временным фильтром «5 лет». (VOSviewer версии 1.6.11). В качестве наиболее перспективных направлений для дальнейших исследований центральных α2-адреномиметиков и их применения в клинической практике определены изучение эффектов и механизмов цитопротекторного и антиоксидантного действия, исследование применения лекарственных средств в моноварианте и в комбинациях для разработки анальгетических препаратов, анестезии и создание комбинированных рецептур с замедленным высвобождением антагонистов с целью нивелирования побочных эффектов.

КЛЮЧЕВЫЕ СЛОВА

Альфа2-адреномиметики, α2-агонисты, дексмедетомидин, комбинации, антиоксидант, цитопrotektion, перекисное окисление липидов.
INTRODUCTION

Medetomidine is a potent and highly specific alpha2-adrenergic receptor agonist (α2-AR), is registered in 34 countries and is widely used abroad. In medico-biological research its dextrorotatory isomer - dexmedetomidine is broadly used [1]. According to its chemical properties, medetomidine is a lipophilic compound that is rapidly and completely absorbed when administered intramuscularly. The half-absorption period is approximately 7 minutes; the peak concentration in plasma is reached in 30 minutes [2]. Dexmedetomidine is a racemic mixture of two stereoisomers, dextro- and levomedetomidine. The dextrorotatory isomer is an active principle of the drug, levorotatory is practically depleted of any pharmacological activity and induces moderate sedation and analgesia only when administered in high doses [3]. The drug can be administered intravenously (iv), intramuscularly (im) and subcutaneously (sc), but the last route of administration is inferior to others in completeness and intensity of sedation [4].

The pharmacological effects of dexmedetomidine, which determine its field of application in medico-biological research, are characteristic of alpha2-adrenergic agonists (α2-AM) and include deep sedation, analgesia, muscle relaxation, anxiolytic effect, the reduction of need for iv and inhalation anesthetics.

Within the pharmacological group, dexmedetomidine is one of the most novel and selective agents, however, it is not devoid of undesirable effects from the cardiovascular system (bradycardia, arrhythmia, hypo- and hypertension, decreased cardiac output), which are observed with the introduction of α2-AM of the first generation (xylazine). This causes some caution among clinicians about the application of dexmedetomidine in premedication and as a sedative agent, especially in a monovariant.

Nevertheless the quantity of publications on this drug is still increasing from 2004 (Fig. 1), and overall number of nonclinical and clinical research in 2018 was 142 and 118 respectively.

The decrease of amount of nonclinical and clinical research of dexmedetomidine in 2019 in the background of preservation of considerable volume of experimental research allows to admit widening of application evidence of a particular drug and representatives of a pharmacological group in general.

Elaboration of experimental data of α2-AM pharmacodynamics as well as the analysis of directions of nonclinical and clinical studies will allow to expand the understanding of possibilities for application of drugs, belonging to this pharmacological group, in clinical practice both in a monovariant and as a part of combined formulations.

Physiological effects of alpha2-adrenergic receptors on the basis of publication analysis of nonclinical safety studies

Adrenoreceptors of α2-type are a separate subclass of alpha2-adrenergic receptors which are localized in the central nervous system (CNS) and in almost all peripheral tissues [5]. Subtypes of α2A, α2B, α2C, α2D are distinguished on basis of differences in their structure, distribution pattern in CNS [6] and sensitivity to pharmacological preparations [5]. Apparent interspecific differences in the ratio of subtypes of α2-AR, their density and localization in CNS are noted.

The most important subtypes of clinical significance can be considered the α2A-subtype, which is responsible for the level of wakefulness in the brain stem, and the α2B-subtype that regulates the diameter of peripheral vessels [7].

The largest role in the implementation of interspecific differences in the effects of α2-AM is assigned to the ratio of receptor subtypes in the brain stem. For example, in the brainstem of dogs and rats α2A-subtype predominates and in the brainstem of sheep it is α2D [8]. It should be noted that ruminants, in view of the features of α2A and α2D subtype functioning in the brainstem, are most sensitive to the action of α2-AM. These features should be taken into account in the prediction of the efficacy and toxicity of α2-AM based on interspecific transfer.

Sedation and analgesia caused by α2-AM depends not only on species characteristics of receptor subtypes, but also on the selectivity of the drug for the α1 or α2 subtype. The majority of well-known α2-AMs are also capable of activating the α1-subtype, this effect makes a significant contribution to the manifestations of pharmacological action, that is especially typical for low-specific agents, such as xylazine. Activation of the α1-subtype causes excitement, anxiety, an increase of locomotor activity and the level of wakefulness [9]. These effects are noted when xylazine is used in high doses: 4-8 mg/kg [10]. It was shown that stimulation of central α1-receptors reduces the hypnotic effect of α2-AM, for example, dexmedetomidine [11]. It was shown that with the introduction of α2-AM in high and toxic doses at the initial period of drug action, the effects of α1-AR activation prevail [12].

When used in therapeutic doses highly selective drugs possess the greatest activity and safety. It is reported that the selectivity ratio between the α2/α1 subtypes of the drugs is the following: medetomidine (1620/1), detomidine (260/1), clonidine (220/1), xylazine (160/1) [13].

Sedative effect

Interest in the usage of α2-AM is stipulated by their ability to cause deep sedation and anxiolytic effect. These

![Figure 1 – The dynamics of publications on dexmedetomidine starting with 2014.](image-url)
effects are mediated by receptors that are predominantly localized in the neurons of the locus coeruleus of the bridge and lower part of the brainstem [10]. Activation of presynaptic α2-AM leads to blockade of the release of norepinephrine necessary to maintain the level of wakefulness. The network effect of impaired release of norepinephrine leads to sedation.

In some cases the impossibility of achieving optimal sedation with the use of α2-AM in therapeutic doses is noted. This may be due to previous stress, fear, agitation, pain and various conditions that contribute to the release of endogenous catecholamines. In clinical practice there are cases when patients were sedated with dexmedetomidine, a temporary “awakening” occurred (a decrease in the degree of depression of consciousness) as a result of exposure to external stimuli [14].

**Analgesia**

α2-AM cause analgesia due to stimulation of receptors at various levels of pain impulse in the spinal cord and brain [15]. Radioligand studies have shown a high concentration of α2-AR in the lateral horns of the spinal cord [16] and in the brainstem, where nociceptive stimuli are processed [17].

In the regulation of pain sensitivity there is an interaction between opioid receptors and α2-AR in the brain [18] and spinal cord [19]. α2-AR and opioid receptors are found in the same areas of the CNS and are often localized at the same neuron. It is believed that both groups of pharmacological drugs produce their analgesic effect according to similar mechanisms related to the activation of the system of secondary messengers through G-proteins, which leads to the opening of potassium channels, hyperpolarization of the postsynaptic membrane and blockade of impulse conduction along the pathways of pain sensitivity.

The results of some experimental and clinical studies indicate that the pronounced analgesic effect of α2-AM is not supported throughout the sedation period. In this regard, drugs of this pharmacological group cannot be used in the monovariant as analgesics for painful or chronic pain have revealed their high activity in systemic and epidural administration [21]. However, the analgesic effect may be accompanied by undesirable sedation and negative effects on the cardiovascular system.

**Effects on the cardiovascular system**

By stimulating the central and peripheral adrenergic receptors, α2-AMs affect the functioning of cardiovascular system. The most pronounced effect is observed in sick and weakened animals, as well as against the background of previous pathology of heart or blood vessels [22]. The main negative effect of α2-AM is bradycardia, bradyarrhythmia (AV blockage of 1-2 degree), a pronounced decrease in cardiac output up to 50%, an increase in total peripheral vascular resistance [23, 24]. According to the results of many studies, it was found that a decrease in cardiac output is not associated with a direct effect of α2-AM on myocardial contractility. It is secondary to an increase of total peripheral vascular resistance and a decrease in heart rate (HR) [25].

As a rule, with the introduction of α2-AM, first an increase in total peripheral vascular resistance occurs, followed by a weakening of the peripheral effect and restoration of the blood pressure (BP) level to normal values [26]. The evidence and duration of an increase in total peripheral vascular resistance and blood pressure during administration of α2-AM depends on a number of factors: selectivity, dose, route of administration (iv or im). The initial hypertonic effect of α2-AM is more pronounced when they are administered in high doses and with intravenous method of administration compared with intramuscular one [27].

According to the analysis of many studies, it can be concluded that when using (dex)medetomidine in therapeutic doses, clinically significant hypotension does not develop. In case of blood pressure was initially elevated, then its value returns to the species norm [28].

**Arrhythmogenicity**

The introduction of α2-AM often causes a decrease in heart rate to 30-50% [29]. Frequent cases of vagal bradycardyrias, AV blockage of 1-2 degree were noted [30]. As a rule, these rhythm disturbances are not life-threatening and are often localized in the neurons of the locus coeruleus of the bridge and lower part of the brainstem [10]. Activation of α2-AR and opioid receptors is associated with a reflex reaction to peripheral vasoconstriction and a decrease in sympathetic tone. With the introduction of α2-AM, an AV block of the 3rd degree and a stop of the sinus node rarely occur [31]. At the same time, in an electrocardiographic study, bradycardia is not accompanied by impaired sinus rhythm [32].

Selective α2-AMs don't produce true arrhythmogenic effect (defined as a decrease in average effective dose of adrenaline for ventricular arrhythmias). In contrast, dexmedetomidine administered intramuscularly causes an increase of the effective dose of adrenaline for ventricular arrhythmias [33].

**Impact on the function of the respiratory system**

Sedation on the background of the introduction of α2-AM is accompanied by a decrease of frequency of respiratory rate (RR). Respiratory depression is secondary to CNS depression, however, when compared with other sedatives, the effect of α2-AM on respiration is not so pronounced even in sublethal doses [9]. In those works where the partial pressure of gases was measured, an increase in pCO2 was noted while maintaining a normal level of pO2 [34].

It has been established that the degree and clinical significance of respiratory depression increases with the use of α2-AM in combination with other sedatives. The
most pronounced respiratory disorders were recorded after the introduction of combinations of medetomidine with opioids and propofol [35]. The combination of medetomidine with ketamine is considered to be much safer [36].

**Muscle relaxation**

α₂-AM in analgesic doses is known to cause muscle relaxation [37]. This effect is associated with inhibition of α₂-AR at the level of the spinal cord interneurons [12]. It should be noted that tizanidine is effective in reducing muscle spasticity in stroke, traumatic brain injury and multiple sclerosis due to the pronounced muscle relaxant effect [9].

**Hypothermia**

With sedation caused by α₂-AM, a decrease in body temperature may occur. In general, a decrease in temperature under the influence of α₂-AM may be associated with inhibition of CNS and a decrease in motor activity [38].

**Fasciculations, cramps**

This reaction develops, as a rule, in a noisy environment, on the basis of which it can be assumed that the mechanism of its occurrence is associated with an increase in sensitivity to sound stimuli [39]. In clinical practice, it is also noted that with dexmedetomidine sedation, “awakening” (increased wakefulness) is possible in patients as a response to external stimulation [40].

**Endocrine manifestations**

The results of various studies indicate that α₂-AM reduces the level of stress hormones during surgery, and thus, can mitigate stress reactions to surgical procedures in dogs [9].

α₂-AMs, especially xylazine, have been reported to increase blood glucose by suppressing insulin release and stimulating glucagon release following activation of beta and alpha pancreatic cells respectively [32]. With respect to selective α₂-AM medetomidine no hyperglycemic effects were detected [20].

Intravenous administration of medetomidine in doses of 10-29 μg/kg induces a diuretic effect lasting up to 4 hours [41].

An increase of the release of growth hormone under the influence of α₂-AM was revealed, however, the clinical significance of this effect has not been conclusively established [3].

**Vomiting**

α₂-AM induces vomiting in animals due to stimulation of chemoreceptors of the trigger zone located in the locus coeruleus region of the brain [42]. Xylazine induces vomiting at the initial period of sedation [43].

According to clinical safety studies of dexmedetomidine in healthy volunteers, vomiting in humans following the administration of the drug develops in 4.6% of cases, depending on the route of administration [4].

**Effect on the digestive tract**

α₂-AM reduces gastric secretion, increases the duration of food transit [9], and inhibits colon motility [44].

**Intraocular pressure**

The multidirectional effect of α₂-AM on pupil diameter and intraocular pressure in animals of various species has been reported [45].

**Intracranial pressure**

α₂-AM reduces central blood flow due to vasoconstriction, and thus reduces intracraniar pressure in dogs anesthetized with isoflurane on mechanical ventilation [46].

The analysis of pharmacodynamics effects of central α₂-AM allows to assume a wide possibility of application of drugs belonging to this group in clinical practice both in a monovariant and in combination with other drugs. Moreover, therapeutic indications for the use of central α₂-AMs are not limited only to use in anesthesiology, which requires a number of additional studies of their effectiveness and safety in the experiment.

**The analysis of trends and directions of research of α₂-am-adrenomimetics with the example of dexmedetomidine**

To determine the trends and directions of research of central α₂-AM the most frequently used drug in clinical practice, dexmedetomidine, was chosen. The analysis used bibliographic data selected in the MEDLINE database.
for the keyword “dexmedetomidine” with a temporary filter of “5 years”. The selected array of information was analyzed using text-mining technology using the graphic visualization method in VOSviewer version 1.6.11 (Center for Science and Technology Research (CWTS) of the University of Leiden, the Netherlands). At the first stage of the study, a conceptual map was constructed based on the frequency of occurrence of keywords and MESH-terms (Figure 2).

The analysis, based on indicators of the relationship between the terms, identified 4 main clusters (in the figure they are presented in different colors). The main directions of research on α2-AM are related to the experimental study of the cytoprotective effect of dexmedetomidine on models of damage of different organs and systems, with clinical safety studies in anesthesiology practice, with application in veterinary medicine. To clarify the results, an in-depth bibliographic analysis of the published materials was carried out, including the full text of the articles (Figure 3).

The presented map made it possible to identify more clearly the main areas of research in the field of α2-AM studies. As the most promising area of research the use of α2-AM cytoprotectors and antioxidants should be considered. The effects of the protective effect on alveolocytes [47], cardiomyocytes [48, 49], hepatocytes [50] are described in details in literature. To date, the mechanisms of such action are at the initial stage of their study. It is known about the reduction of lipid peroxidation products, the production of peroxynitrile radicals [51], the stimulation of the activity of endogenous glutathione and superoxide dismutase [52]. A significant role in this case is given to stabilization of the inner mitochondrial membranes [53, 54]. In our opinion, under the conditions of modeling damage of various types, accompanied by depletion of the antioxidant systems of the body, an increase in the effectiveness of dexmedetomidine can be achieved by restoring the pool of water-soluble cytosolic antioxidants.

Another important block of research is the study of the combined effect of α2-AM with drugs of other pharmacological groups [55], which to a greater extent potentiate the analgesic or anesthetic effect of each other. These studies are naturally associated with clinical trials with various groups of people [56, 57, 58].

Separate studies are devoted to the research of combined application of agonists and antagonists of α2-AM [55]. In these publications, the authors set themselves a goal to stop possible side and undesirable effects, as well as to achieve more controlled sedation.

Based on the analysis, we can formulate several of the most promising trends and directions in the study of central α2-AM, which determine the expansion of indications for their future application in medico-biological research:

1. The study of the effects and mechanisms of cytoprotective and antioxidant effects, including their combined administration with non-enzymatic antioxidants. The effects associated with stabilization of the inner mitochondrial membrane are of particular interest.

2. The study of combinations of painkillers intended for application in self-help and mutual assistance, as well as first aid.

3. The study of its own anesthetic effect in a monovariant. In this case, it will be extremely important to conduct in-depth experiments to assess safety during the administration of drugs in sublethal doses, as well as to assess changes in the functioning of external respiration and indicators of the cardiovascular system.

4. The study of drug combinations containing central α2-AM antagonists (for example, atipamezole) with delayed controlled release in order to stop possible side and undesirable effects.

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