Effect of Sigma Receptor Ligand Haloperidol in Guinea Pig Heart

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Abstract

Sigma receptors represent a group of enigmatic binding sites with many functions in different tissues. In rat heart, they are responsible for modulation of cardiac contractility. They are also responsible for some adverse side effects of certain psychotropic drugs on cardiovascular system. In this study, the effect of sigma receptor ligand haloperidol on guinea pig heart multicellular preparations was studied. Its positive inotropic effect previously described in rat cardiomyocytes was confirmed in left atria and papillary muscles isolated from the right ventricles (an increase of the twitch amplitude by 36% to 123% in comparison with the control period in atria and by 9% to 46% in ventricular preparations; all values were statistically significant when compared to control in atrial preparations, but only in the first five min of perfusion in papillary muscles). Also desensitisation process previously reported in rat myocardium after repeated exposure to haloperidol was confirmed (although not statistically significant). The normalized course of mechanical restitution curve was not changed during perfusion with haloperidol. It is concluded that its positive inotropic effect is very likely mediated via increased cytosolic availability of calcium.

Cardiac sigma receptor, contractility, mechanical restitution, desensitisation

Psychotropic drugs quite often exert marked cardiac side effects (increase in blood pressure, arrhythmias, palpitations and even sudden cardiac death), mainly mediated by central and autonomic nervous systems. Some of them are, however, caused also by direct effect of different compounds on cardiac tissue. Sigma receptors are among the binding sites responsible for these direct effects and many sigma ligands belong to different groups of drugs used in everyday clinical practice for management of pain, agitated delirium, treatment of psychoses, etc.

The sigma receptors were first reported in central nervous system. Initially, they were considered a mere subtype of the opioid receptor family, but later they were conclusively defined by binding studies as distinct from any other known class of receptors. Since, they have been found also in many non-neural, peripheral tissues (immune system, digestive tract, endocrine and reproductive systems and excretory system, myocardium, etc.) in various species, including humans. They are present in many tumours and metastases in high densities. Three subtypes of these receptors are reported – sigma1, sigma2 and sigma3 (reviewed by Nováková 1998).

Although sigma receptor has been repeatedly cloned in several tissues and species (Hanner et al. 1996; Kekuda et al. 1996; Seth et al. 1998), identification of its endogenous ligand has not yet been successful. Among many candidates, progesterone takes a prominent place.

In cardiac tissue, modulation of contractility by sigma receptor ligands was first reported in rat neonatal cultured cardiomyocytes (Elia et al. 1994). Later, sigma receptors were found in the membranes of adult rat ventricular cardiomyocytes (Nováková et al.
1995). Also data on the effects of sigma ligands in rat isolated hearts have been reported (Maslov et al. 1997, 1999; Nováková et al. 2001) and on desensitisation of sigma receptors in heart muscle by repeated treatment with sigma ligands (Nováková et al. 2001; Ela et al. 1996).

The cardiac effects of sigma ligands thus have been studied so far exclusively on rat heart, which is well-known to be rather exceptional among the mammalian species. The aim of this study was to investigate the effect of haloperidol in guinea pig multicellular heart preparations. This model was chosen since this experimental set-up is reliable and the experimental conditions can be strictly controlled.

Materials and Methods

In the study 19 left atria and 15 papillary muscles from the right ventricle of male guinea pigs were included. The body weight of animals varied from 230 to 360 (average, 295 ± 28) grams. The experiments were restricted to males to exclude possible effect of progesterone on female myocardium.

The animals under deep ether anaesthesia were sacrificed by cervical dislocation. The chest was quickly opened, the heart immediately removed and placed in a preparation bowl with a cold (5 °C) Krebs-Henseleit solution of the following composition: NaCl, 118 mM; NaHCO₃, 24 mM; KCl, 4.2 mM; KH₂PO₄, 1.2 mM; MgCl₂, 1.2 mM; CaCl₂, 1.2 mM; and glucose, 5.5 mM. The right ventricle was opened and all suitable papillary muscles removed. Then, the left auricle strip was cut off and placed together with the right ventricle papillary muscle in the perfusion bath (Fig. 1).

We employed a horizontal, plastic, double-walled, thermostatically controlled bath for pharmacological studies. It contains a pair of stimulation electrodes at the bottom, one for stimulation (under the clips that held the preparations), the other for grounding. The bath was filled with 10 ml Krebs-Henseleit solution aerated with mixture of O₂ and CO₂ (95:5%). Both muscles were fixed by a clip at one end and attached to a mechano-electric transducer by a thread bound to its opposite end. Simultaneous recording of the tension generated by the preparations was performed under isometric conditions.

The signals were amplified and digitised at sampling rate of 250 Hz by a one-channel bipolar 16-bit A/D converter. The specialised software allowed simultaneous 30-min record of two trends - maximal and minimal tension of the preparations after stimulus and up to ten 15-second snap-shots with detailed (sampling period of 4 ms) recording of stimulated twitches. The recorded data were provided with identification labels, saved in archive files, exported into ASCII and worked out in Excel.

The effects of specific sigma receptor ligand haloperidol on the amplitude of twitch and on the restitution of contractility in both atrial and ventricular guinea pig heart preparations were investigated. The experiments were carried out at 30 °C and fluctuations in temperature did not exceed 0.5 °C. The preparations were stimulated with 1ms pulses of twice the current threshold. Basic stimulation frequency was 1 Hz throughout the experiments.

Fig. 1. Horizontal perfusion bath for heart multicellular preparations (A, side view; B, from above). 1. stimulation electrodes; 2. thermometer; 3. grounding electrode; 4. bubbling; 5. tensometers; 6. clips.
Initially, the preparations were stimulated for 30–45 min. At the end of this period control contractions were recorded. Then the contractions of variable prematurity were recorded according to the protocol (Fig. 2) and the mechanical restitution curve was constructed. Next, sigma receptor ligand haloperidol at a concentration 10 nM was administered to the bath for 30 min. Snap-shots of 15 contractions at basal rate (1 Hz) were recorded after 1, 3, 5, 10, 15, 20, 25 and 30 min. Then, the stimulation protocol for assessment of drug affected restitution curve was repeated. This protocol was employed once again after 15 min washout with Krebs-Henseleit solution. Eventually, haloperidol at the same concentration was administered to the bath for the second time in order to test the possible desensitisation.

Obtained data were tested for normality by the means of Kolmogorov-Smirnov test. Statistical significance was assessed using one-sample $t$-test, two sample $t$-test or analysis of variance (ANOVA). All statistical analyses were two-sided and performed at the 5% significance level.

**Results**

We observed a marked positive inotropic effect previously described in rat ventricular cardiomyocytes in all experiments. In atria, an increase of the twitch amplitude by 36% to 123% in comparison with the control period (average of control twitches amplitudes represents 100%), with the maximum effect lasting for the first 5 min of perfusion. All values were statistically significant when compared to control. In ventricular preparations, the increase of contraction by 9% to 46% peaked during the first minute of perfusion (Fig. 3). The increase was statistically significant (compared to control values) only in the first five min of perfusion with haloperidol.

The second administration of haloperidol caused a positive inotropic effect in both preparations again. In atrial preparations, however, this effect was attenuated in the whole course of perfusion (increase of contractions by 9% to 94% in comparison with control period, with a maximum effect in the third minute of perfusion), (Fig. 4). When

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**Fig. 2.** The experimental protocol for construction the mechanical restitution curve. A, B – last two steady-state contractions in a train of regular stimuli. C – contractions in response to single stimuli of variable prematurity. The amplitudes of the successive premature contractions (C) form the restitution curve.

**Fig. 3.** Time course of the effect of sigma ligand haloperidol on normalized contraction amplitudes of guinea pig atria and papillary muscles. Temperature, 30 °C; stimulation frequency, 1 Hz. Mean ± SD of 15 experiments. Wash-out not shown.
comparing the first and the second haloperidol administration, no significant difference was found.

In right ventricular papillary muscles, the positive inotropic effect was attenuated more clearly during the second administration of haloperidol – the increase of contractions ranged from 9 to 38%, with the maximum between the 10th and the 30th minute of perfusion), (Fig. 5). Again, no significant difference between the first and the second haloperidol administration was found.

Fig. 4. The effect of two administrations of haloperidol (10 nM) on contractions of guinea pig left atria. Temperature, 30 °C; stimulation frequency, 1 Hz. Normalized mean ± SD from 19 experiments. Wash-out not shown.

Fig. 5. The effect of haloperidol (10 nM) on right heart papillary muscles. For legend see Fig. 4. Normalized mean ± SD from 15 experiments.
The pattern of mechanical restitution was not altered in either preparation under the effect of haloperidol (Figs 6, 7). There were no significant differences between mechanical restitution curves in either situation or preparation.

Fig. 6. The normalized mechanical restitution of left atrial muscles under control conditions, after 30 min of perfusion with haloperidol and after 15 min of wash-out. Abscissa, extrainterval, period between a regular and a premature stimulation pulse. Temperature, 30 °C; basal stimulation frequency, 1 Hz. Mean ± SD from 19 experiments.

Fig. 7. Mechanical restitution of right heart papillary muscles. For legend see Fig. 6. Normalized mean ± SD from 15 experiments.
Discussion

In this study, the effect of haloperidol, a typical sigma receptor ligand was investigated in guinea pig heart multicellular preparations. Haloperidol is a psychotropic drug used in everyday clinical practice for treatment of psychoses like schizophrenia or severe agitated delirium. Its cardiac side effects - arrhythmias such as torsade de pointes, ventricular fibrillation and other life-threatening complications and even cardiac arrest - are well known and were reported repeatedly (Di Salvo and O’Gara 1995; Cruz et al. 1983; Tisdale et al. 1991; Lawrence and Nasraway 1997). The concentration used in our experiments was 10 nM. It was chosen on the basis of our previous experiments (Nováková et al. 1995) where binding studies revealed that $K_i$ for sigma receptors in rat cardiac myocytes is $6.1 \pm 1.3$ nM. Since many drugs in clinical use bind to sigma receptors with nanomolar affinities (certain neuroleptics, monoamine oxidase inhibitors and several antihistamines – Vilner et al. 1995), we consider this concentration relevant.

Since the so far data have been obtained in myocytes and in preparations from rat heart which is quite exceptional from many points of view, the effect of sigma binding on the contractility of guinea pig heart multicellular preparations was examined. Inotropic effect was first described in neonatal rat cultured cardiomyocytes (Elia et al. 1994): exposure to nanomolar concentrations of the sigma receptor ligands (+)-3-PPP ((+)-3-hydroxyphenyl-N-(1-propyl)piperidine), (+)-pentazocine and haloperidol, induced specific pattern of changes in twitch amplitude of the cardiomyocytes, consisting of an initial decrease, followed by a transient increase and a final decrease of amplitude.

In similar experiments with adult rat isolated ventricular myocytes, the initial decrease of the amplitude of twitch was absent and the increase in the amplitude was much higher in most of the cells (Nováková et al. 1995). In part of the cells exposed to low concentrations of sigma receptor ligands, only a decrease of the contractile response was observed. In analogous experiments carried out with highly selective, high affinity sigma receptor ligands BD-737 (1S, 2R-cis-N-[2-(3,4-dichlorophenyl)ethyl]-N-methyl-2-(1-pyrrolidinyl) – cyclohexylamine) and BD-1047 (N-[2-(3,4-dichlorophenyl)ethyl]-N,N’,N’ – trimethylethylenediamine) the positive inotropic effect was also observed (Nováková et al. 1998). All these data were obtained with nanomolar concentrations of sigma ligands. These experiments brought evidence in support of the idea that the cardiotropic effects of psychotropic drugs used are due to binding to specific receptors and not by a cross-talk between the different receptor systems.

In all the above mentioned experiments fluctuations of the response to sigma ligands was present. This typical pattern can be observed also in the time course of the effect of haloperidol in guinea pig heart preparations (Fig. 3).

Recently, modulation of the negative inotropic effect of haloperidol by drugs with positive inotropic effects in isolated rabbit heart has been reported (Hatip-Al-Khatib and Bolukbasi-Hatip 2002). However, the negative inotropic effect was elicited by micromolar concentration of haloperidol, which is by three orders of magnitude higher than the concentration used in our study. In such case, other receptor systems may play a role in this effect (e.g. dopaminergic receptors).

Desensitisation of sigma receptors was reported in neonatal cultured rat cardiomyocytes (Elia et al. 1996) and in neurons (Inoue et al. 2000). In isolated rat hearts perfused according to Langendorff, the exposure to haloperidol caused prominent ventricular ectopic activity and bigeminy in all hearts. The occurrence of arrhythmias was significantly lower during the second haloperidol administration in support of the concept of desensitisation (Nováková et al. 2001). In the present study, repeated treatment of guinea pig heart preparations with haloperidol seems to trigger similar desensitisation process as described previously in rat heart muscle. However, attenuation of positive inotropic effect caused by
the second haloperidol administration was statistically not significant. Thus, the desensitisation of sigma receptors in guinea pig heart deserves more investigation.

The mechanical restitution curve is a sensitive tool for estimation of heart preparation viability. Its course is not changed under the effect of haloperidol and does not change even during long lasting desensitisation experiments. We can assume that positive inotropic effect of sigma ligand is mediated via increased cytoplasmic Ca\textsuperscript{2+} concentration as previously reported in rat myocardium.

Thus, the positive inotropic effect of sigma receptor ligand haloperidol was confirmed also in guinea pig atrial and right ventricular heart preparations. Attenuation of this positive inotropic effect of haloperidol in the case of repeated exposure leads to the conclusion that sigma receptors in guinea pig heart probably undergo a desensitisation process previously reported in rat myocardium.

Úãinek sigma ligandu haloperidolu na srdce morãete

Sigma receptory pfiedstavují skupinu ménû prozkouman˘ch vazebn˘ch míst s rozmanit˘mi funkcemi v mnoha tkáních. U srdce potkana se podílejí na jemné modulaci kontraktility. Jsou také odpovûdn˘ za neÏádoucí kardiovaskulární vedlejš˘ úãinky nûkter˘ch psychotropních látek. V´ této práci byl zkoumán úãinek ligandu sigma receptorÛ haloperidolu na multicelulární preparáty ze srdce morãete. Pozitivní inotropní úãinek této látky popsan˘ dûrve u kardiomycytÛ potkana byl potvrzen i u lev˘ch síní a papilárních svalÛ izolovan˘ch z prav˘ch komor (nárûst amplitudy stahu mezi 36% a 123% oproti kontrole u síní a mezi 9% a 46% u komorov˘ch preparátÛ; v‰echny hodnoty statisticky v˘znamné u síní, ale pouze v prvních 5 minutách perfuze u papilárních svalÛ). ZároveÀ byla potvrzena i desensitizace sigma receptorÛ, která byla popisována u srdce potkana po opakované aplikaci haloperidolu (statisticky nesignifikantní). Normalizovaná kﬁivka mechanické restituce se pod vlivem haloperidolu nemûní. Pozitivnû inotropní úãinek haloperidolu je pravdûpodobnû zprostfiedkován zv˘‰ením cytosolické dostupnosti vápníku.

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