Effects of the NO donor sodium nitroprusside on oxygen consumption and energetics in rabbit myocardium

Abstract Nitric oxide (NO) has influence on various cellular functions. Little is known of the influence of NO on myocardial energetics. In the present study oxygen consumption and mechanical parameters of isometrically contracting rabbit papillary muscles (1 Hz stimulation frequency) were investigated at varying interventions while maintaining physiological conditions (37°C; 2.5 mM Ca2+) to study the effects of NO on energetics. The NO donor sodium nitroprusside (SNP) showed a negative inotropic effect. SNP decreased the maximal force in normal rabbit muscle strips by 30%, the force time integral (FTI) by 40% and the relaxation time by 20%. In addition the oxygen consumption decreased by 60%, a notably disproportional decrease compared to the mechanical parameters. Consequently, the economy as a ratio of FTI and oxygen consumption is significantly increased by SNP. In contrast the negative inotropic effect due to a reduction in extracellular Calcium (Ca2+) from 2.5 to 1.25 mM reduced FTI and oxygen consumption proportionally by 40% and did not change economy. The effect of NO on force and oxygen consumption could be reproduced by the application of the cyclic guanosine monophosphate (cGMP) analogue 8-bromo-cGMP. In summary, NO increased the economy of isometrically contracting papillary muscles. The improvement in contraction economy under NO seems to be mediated by cGMP as the secondary messenger and maybe due to alterations of the crossbridge cycle.

Key words energetics – NO – myocardium – oxygen consumption

Introduction

Nitric oxide (NO) has been shown to be of significance and relevance for short and long term regulation of myocardial function and gene expression.

The first evidence that NO may influence myocyte function resulted from observations of changes in myocardial contractility by the release of various transmitters of the endocard or endothelium [41, 42]. NO and other substances which increase the intracellular cGMP level, decreased the systolic tension in single myocytes, muscle strips and isolated hearts. Simultaneously, single contraction parameters, especially contraction time, were also altered. The acceleration in relaxation time was the main reason for the significant shortening of the contraction time [20, 35, 37, 42].
Both, basal and cytokine induced NO synthesis influence the effects of β-adrenergic stimulation. Several experiments have shown that the positive inotropic effect of isoprenaline was significantly decreased in the presence of NO [1, 39, 46, 47]. This effect is nearly eliminated by the inhibition of NO-synthases through L-NMMA.

The influence of NO on myocardial inotropic, chronotropic and dromotropic response seems to be biphasic [36]. Kojda et al. [15] observed that isolated ventricular rat myocytes exhibited a small, however significant positive inotropic reaction as well as an elevated isoprenaline effect under a small concentration of the NO-donor S-Nitroso-N-Acetyl Penicillamine (SNAP). In the same experiment higher doses of SNAP decreased maximal systolic tension.

The endocardium plays an important role in the regulation of the myocardial function by NO. Fort et al. [5] showed that the intraventricular infusion of SNP at any concentration into hearts without endocardium had no effect on maximal tension and relaxation time. However, if SNP was added to coronary arteries with endothelium, the left ventricular tension decreased and the relaxation time was shortened. Moreover, in papillary muscles isolated from pigs with the endocardium removed, SNP showed a positive inotropic outcome. These findings validate the influence of endocardium or respectively endothelium on the decrease in systolic tension and shortened relaxation time in myocardium.

The observed changes in muscle mechanics led to the supposition that NO may influence the oxygen consumption as well as energetics of myocardium. To identify the effects of NO on energetics we investigated the influence of the NO donor SNP on papillary muscle contractions and oxygen consumption. The effects of NO were compared with the effects of cGMP as well as with a change in extracellular Ca2+ concentration.

Methods

Papillary muscle preparation

Animal handling was reviewed and approved by the Animal Investigation Committee of the University of Goettingen. Right ventricular papillary muscles were gathered from New Zealand White rabbits (2.2–2.8 kg body weight). Immediately after cardiectomy, the heart was removed and submerged in cold (4°C) protective solution bubbled with 95% O2 and 5% CO2 (Carbogen). The solution contained (mmol/l) Na+ 152, K+ 3.6, Cl− 135, HCO3− 25, Mg2+ 0.6, H2PO4− 1.3, SO4 2− 0.6, Ca2+ 2.5, glucose 11.2 and 2,3-butanedione monoxime (BDM) 30 along with 10 IU/l insulin (Hoechst, Inc). All chemicals were purchased from Sigma (Sigma, Inc) unless otherwise indicated. This protective solution was shown to preserve myocardium during transportation and dissection and to be completely reversible after washout [24]. Ventricles were gently rinsed to clear remaining blood. Preparation of myocardium was performed in a dissection chamber filled with the protective solution. Right ventricular papillary muscles were abscessed at the base from the ventricular wall. Only long, cylindrical muscles were used.

Experimental protocol

Papillary muscles were placed into the measurement apparatus (Muscle Research System, Scientific Instruments, Heidelberg, Germany) and clamped at both ends. The muscles were perfused with Krebs-Ringer solution (composition as above without BDM) bubbled with Carbogen and warmed to 37°C. Hepes Buffer with a concentration of 5 mmol/l at a pH of 7.4 was used in addition to the Krebs solution during the SNP experiments. Isometric contraction was accomplished by stimulation with a 5 ms pulse applied end-to-end at a voltage 25% above threshold and a frequency of 1 Hz. After an equilibration period of 30 min, muscles were stretched in 0.05 mm increments until additional stretch did not produce an increase in developed force. This length was taken as lmax. Mechanical parameters (twitch tension and timing parameters) and oxygen consumption were measured at steady state conditions. Dose–response experiments were performed with incremental doses of SNP and 8-bromo cGMP in each muscle preparation.

Isometric force was acquired digitally from the force transducer (KG4, Scientific Instruments). Twitch tension was defined as the active tension developed during the isometric twitch. Force-time integral was calculated as the area under the twitch force curve. At the end of each experiment muscle length and weight were measured. Cross-sectional area for normalization of force values was calculated as the ratio of blotted muscle weight to muscle length (lmax).

Oxygen consumption

The method used to measure oxygen consumption was described and validated previously [21]. Briefly, a miniature oxygen electrode continuously measured PO2 in the experimental chamber close to the muscle surface. The drop in PO2 at the probe tip was measured over 20 s and compared to precalculated theoretical profiles to determine the rate of oxygen
consumption of the muscle (mVO₂). Economy was calculated at each concentration of SNP and cGMP as the ratio of FTI to oxygen consumption.

Statistics

All results are reported as means ± SD unless otherwise noted. Comparisons between NO treated and control muscles were performed by unpaired t test or one-way ANOVA with a Tukey adjustment for multiple comparisons. A P value < 0.05 was considered significant for all tests.

Results

Dose–response experiments in 8 isometrically contracting papillary muscles from different animals were performed at 10⁻⁵, 10⁻⁴, and 10⁻³ M SNP. Active tension and FTI both decreased continuously with maximum depressions of 29 ± 3% and 36 ± 3% respectively at 10⁻³ M SNP (P < 0.01) compared to control conditions at 2.5 mM free calcium (Fig. 1) while diastolic tension remained unchanged. Consistent with previous experiments [27] SNP significantly decreased the time from peak twitch tension to 90% relaxation (RT90) by 20 ± 5% (P < 0.01). In addition, maximal relaxation velocity (−dT/dt) increased by 20 ± 7% (P < 0.05). Time to peak tension (TTPT) and maximal velocity of force rise (+dT/dt) were not found to be significantly altered (Fig. 2).

The free calcium concentration of the Krebs–Ringer solution was halved from 2.5 to 1.25 mmol/l to compare the observed negative inotropic effects of SNP. This intervention resulted in a 50 ± 3% decrease in active force (P < 0.01) and a 37 ± 6% decrease in FTI (P < 0.01) (Fig. 1). Time to peak tension, +dT/dt and relaxation parameters were not affected (Fig. 2).

Muscle preparations showed a concentration dependent decrease in oxygen consumption with maximal reduction of 55 ± 4% being reached at 10⁻³ M SNP (P < 0.01). Halving the free calcium concentration resulted in a 40 ± 6% decrease of oxygen consumption (P < 0.01) (Fig. 3). Though basal oxygen consumption under SNP and calcium did not differ (P = NS).

Economy as a ratio of FTI to oxygen consumption increased under rising SNP concentrations up to 140 ± 9% (SNP 10⁻³ M; P < 0.01). However, the change in free calcium concentration did not show a significant difference in economy (P = NS) (Fig. 4).

To investigate the possible role of increased intracellular cGMP levels through SNP, mechanical parameters and oxygen consumption were measured in another eight papillary muscle preparations.

Increasing concentrations of 8-bromo cGMP (10⁻⁶, 10⁻⁵ and 10⁻⁴ M) were added to the Krebs–Ringer solution. Twitch force and FTI decreased continuously with increasing cGMP concentrations. At 10⁻⁴ M 8-bromo cGMP twitch force and FTI reached were mini-
with reduction by 28 ± 6% and 37 ± 8% respectively (both \( P < 0.01 \)). Consistent with SNP experiments RT90 was reduced by 19 ± 4% and \( \Delta T/\Delta t \) increased by 26 ± 7% (both \( P < 0.05 \)). TTPT and \( \Delta T/\Delta t \) were not affected. Halving of free calcium in the Krebs–Ringer solution showed similar results as described above (twitch force –47 ± 4%, FTI –42 ± 6%, no change in TTPT and relaxation parameters).

Oxygen consumption of cGMP treated muscle preparations decreased by 56 ± 3% (\( P < 0.005 \)) whereas the halving of calcium resulted in a 43 ± 5% reduction (\( P < 0.005 \)) (Fig. 3). Therefore, the economy for cGMP treated muscles increased by 40 ± 8% (\( P < 0.01 \)) while the change in calcium concentration had no influence on economy (\( P = \text{NS} \)) (Fig. 4).

Discussion

In the present study SNP had a negative inotropic effect in isolated papillary muscles. These results are consistent with studies done in guinea pigs [2], cats [22, 23] and clinical studies with intracoronary infusion of SNP in humans [27]. The additionally described shortening of contraction time and the acceleration of relaxation could also be reproduced [7, 27]. However, some authors [2] observed no significant changes of the relaxation parameters or contraction time in guinea pig hearts. Species and tissue differences could be responsible for these differences.

The negative inotropic effect of SNP was associated with a disproportionate decrease in oxygen consumption as compared to the proportional decrease in active force and oxygen consumption induced by reducing the \( Ca^{2+} \) concentration from 2.5 to 1.25 mM. As a consequence contraction economy increased with SNP whereas halving the calcium concentration had no influence on myocardial energetics.

The participation of NO in the ischemic preconditioning cell-survival program is widely discussed in the literature. Previous studies [17, 18, 40] indicated that endothelium-derived NO is an important regulator of tissue oxygen consumption in myocardium in vivo and vitro. The mechanisms underlying cytoprotection involve the modulation of mitochondrial function such as reversal inhibition of cytochrome c oxidase and a reduction of mitochondrial enzyme activities including aconitase in the Krebs cycle and complex I and II of the mitochondrial electron transport chain.

Peroxinitrate, formed by the interaction of NO with superoxide, also promotes a suppression of respiration [43]. However, the inhibition of cellular respiration by peroxinitrate is not reversible. Thus, it is unlikely to act as a potent physiological regulator for tissue oxygen consumption.

Additionally, Rickover et al. [34] described a SNP triggered activation of the sarcoplasmic calcium-ATPase with increased \( Ca^{2+} \) uptake into the sarcoplasmic reticulum and thus prevention of cytosolic \( Ca^{2+} \) overload in rat cardiomyocytes. The overproportional decrease of oxygen consumption and increase in contraction economy after SNP treatment observed in our study may result from effects on calcium cycling and be part of the beneficial effects of NO during ischemia/reperfusion when oxygen is limiting and tissue becomes acidotic and may enhance ischemic tolerance during preconditioning.

Moreover, NO has been shown to control substrate utilization of myocytes [33, 45]. Lack of NO causes preferential utilization of carbohydrates by myocardium. This deficit may be important especially in heart failure and diabetes when NO production by blood vessels is reduced or abolished. Recchia [32] hypothesized that prevalent carbohydrate utilization might be advantageous for the heart because the ATP yield from carbohydrate oxidation is greater for a
given rate of oxygen consumption, due to the higher ATP/oxygen ratio compared to that of free fatty acid oxidation. Switching from free fatty acid to lactate can result in a reduction of around 10% of oxygen consumption [30]. However, healthy hearts in the fasting state consume primarily fatty acids, because this is their optimal substrate. The shift to prevalent carbohydrate utilization by the failing heart likely represents an adaptive response to face profoundly altered mechanics and energy turnover e.g. increased basal oxygen consumption and reduced efficiency, even if it could be disadvantageous in the long-term.

To further investigate the role of NO on subcellular mechanisms in myocytes, the membrane permeable cGMP-analogue 8-bromo cGMP was used. Through experimentation it was determined that 8-bromo cGMP produced similar results to SNP at only one tenth of the SNP concentration. These findings suggest that the effects of NO on myocytes are primarily mediated by cGMP and consistent to recent studies [16].

Guanylate cyclase is stimulated by extracellular NO and synthesizes cGMP from guanosine triphosphate (GTP). Within the myocardial cells cGMP and cyclic adenosine monophosphate (cAMP) are antagonists. Previous studies described dose dependent effects on contractile function of cGMP or rather NO [15, 31]. On one hand very low NO concentrations increase intracellular cAMP levels by activation of adenylyl cyclases directly [14] and by a cGMP-mediated inhibition of phosphodiesterase [29, 35] with a consequently diminished breakdown of cAMP. This leads to phosphorylation and thus activation of the L-type calcium channel and other key proteins in calcium cycling [38, 44]. Therefore, calcium influx and calcium transients are increased in myocytes resulting in an increase in contractile function but also myocardial oxygen consumption. Moreover, positive inotropic effects of low-dose NO are possible by increasing contractile calcium responsiveness [10]. On the other hand high NO concentrations activate a cGMP dependent cAMP phosphodiesterase with consequent reduction in cAMP concentrations [4, 35]. Additionally, the cAMP phosphorylated L-type calcium channel is dephosphorylated by cGMP dependent phosphatase(s) [38] and troponin I and phospholamban are phosphorylated [13, 16]. Phosphorylation of troponin I reduces the calcium sensitivity of the thin filament by a decrease of the affinity of troponin C to calcium and amplifies the loss in tension [12, 16, 25]. Evidence that high cGMP concentrations cause a negative inotropic effect was shown by extracellular application of acetylcholine [6]. In other experiments with similar results a cGMP-analogue was used [35]. Moreover, phosphorylated troponin I prevents calcium-independent diastolic cross-bridge cycling [26]. Phosphorylation of phospholamban activates the sarcoplasmic calcium-ATPase and thereby accelerates calcium elimination out of the cytosol [28]. A further decrease in systolic and diastolic force and additional shortening of contraction time would result. The described effects are in agreement with our findings of a shortened relaxation time and an increase in relaxation velocity. However, consistent with previous experiments [11] there was no significant change in diastolic force generation assuming that diastolic stiffness is negligible in nonfailing rabbit myocardium in contrast to patients with dilated cardiomyopathy or aortic stenosis.

Thus, the negative inotropic effect of SNP observed in our experiments may be caused by dephosphorylation of the L-type calcium channel, a decreased cAMP concentration in the cell, an accelerated calcium uptake in the sarcoplasmic reticulum or a shift in thin filament calcium sensitivity.

The mechanism behind the improvement of the contraction economy is not clear. Economy improvement may be due to the increases in economy of ATP synthesis or a reduction of ATP usage per unit of developed tension. Animal models have shown that under special experimental conditions NO reduces the ATP synthesis in mitochondria [3]. However, basal oxygen consumption under NO and control conditions did not differ in our experiments, suggesting the economy of ATP synthesis remained unchanged (data not shown). Due to technical reasons high solution oxygen partial pressures are needed in the diffusion method. Therefore, effects of super-physiological oxygen partial pressures on mitochondria NO interactions and NO metabolism can not be excluded and may have an impact on the observed results.

A reduction of ATP usage per unit of developed tension may be a result of changes in excitation-contracting coupling or myofibrillar performance. Both require energy during the contraction-relaxation cycle [19]. Given that excitation-contracting coupling requires 15–20% of the total ATP used [8], it is unlikely that the demonstrated improvement in contraction economy is based on effects on excitation-contracting coupling alone. Moreover, decreased calcium sensitivity by phosphorylation of troponin I may rather increase energy consumption for calcium cycling [12, 25]. Therefore, a major reduction of ATP usage seems to be caused by improved economy at the level of myofibrillar proteins. The underlying mechanism may be an increase in the FTI of a single cross-bridge through a lengthened thin filament attachment time to myosin. A corresponding change of cross-bridge economy was demonstrated with some calcium sensitizers [9] as well as in hypertrophic and insufficient myocardium [8]. This effect caused by NO is most likely responsible for the improvement in contraction economy.
The more pronounced decrease in oxygen consumption relative to FTI and therefore the increase in economy of isolated papillary muscle preparations after SNP treatment may be part of the improvement observed in nitrate treated patients with coronary heart disease.

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