Activation of Distinct cAMP-Dependent and cGMP-Dependent Pathways by Nitric Oxide in Cardiac Myocytes

Martin G. Vila-Petroff, Antoine Younes, Josephine Egan, Edward G. Lakatta, Steven J. Sollott

Abstract—Nitric oxide (NO) donors were recently shown to produce biphasic contractile effects in cardiac tissue, with augmentation at low NO levels and depression at high NO levels. We examined the subcellular mechanisms involved in the opposing effects of NO on cardiac contraction and investigated whether NO modulates contraction exclusively via guanylyl cyclase (GC) activation or whether some contribution occurs via cGMP/PKG-independent mechanisms, in indo 1–loaded adult cardiac myocytes. Whereas a high concentration of the NO donor S-nitroso-N-acetylpenicillamine (SNAP, 100 μmol/L) significantly attenuated contraction amplitude by 24.4 ± 4.5% (without changing the Ca2⁺ transient or total cAMP), a low concentration of SNAP (1 μmol/L) significantly increased contraction amplitude (38 ± 10%), Ca2⁺ transient (26 ± 10%), and cAMP levels (from 6.2 to 8.5 pmol/mg of protein). The negative contractile response of 100 μmol/L SNAP was completely abolished in the presence of the specific blocker of PKG KT 5823 (1 μmol/L); the positive contractile response of 1 μmol/L SNAP persisted, despite the presence of the selective inhibitor of GC 1H-[1,2,4]oxadiazolo[4,3-a]quinoxalin-1-one (ODQ, 10 μmol/L) alone, but was completely abolished in the presence of ODQ plus the specific inhibitory cAMP analog Rp-8-CPT-cAMPS (100 μmol/L), as well as by the NO scavenger oxyhemoglobin. Parallel experiments in cell suspensions showed significant increases in adenylyl cyclase (AC) activity at low concentrations (0.1 to 1 μmol/L) of SNAP (AC, 18% to 20% above basal activity). We conclude that NO can regulate both AC and GC in cardiac myocytes. High levels of NO induce large increases in cGMP and a negative inotropic effect mediated by a PKG-dependent reduction in myofilament responsiveness to Ca2⁺. Low levels of NO increase cAMP, at least in part, by a novel cGMP-independent activation of AC and induce a positive contractile response. (Circ Res 1999;84:1020-1031.)

Key Words: nitric oxide signaling ■ cGMP ■ cAMP ■ contractility ■ cardiac myocyte

Nitric oxide (NO) has been implicated as a mediator of many cellular processes, including endothelium-dependent relaxation of blood vessels, chemical communication between peripheral nerves and smooth muscle, inhibition of platelet aggregation, immune responses, and neurotransmission. These effects of NO have been ascribed to the activation of soluble guanylyl cyclase (GC), leading to the production of cGMP and activation of cGMP-dependent protein kinase (PKG). Furthermore, NO and NO donors have been shown to elicit a wide range of effects on myocardial contractility, but the specific nature of the subcellular mechanisms underlying the diverse effects in heart is largely unknown.

Although the coronary endothelium is responsible for the bulk of the endogenous, physiological production of NO in the heart, NO can also be produced within the cardiac myocytes themselves (ie, autocrine production) by the constitutive NO synthase NOS-3. Dynamic regulation of NO production in the heart apparently yields beat-to-beat oscillations in response to changes in coronary flow and myocardial loading and achieves micromolar levels in close vicinity to the cardiac myocyte. Until recently, it was generally accepted that NO induced negative inotropic effects in cardiac preparations, mediated principally through cGMP-related mechanisms, specifically via the reduction of Ca2⁺ influx through L-type Ca2⁺ channels, either through activation of cGMP-dependent phosphodiesterase or PKG and/or phosphatases. cGMP has also been shown to decrease relative myofilament response to Ca2⁺ and therefore enhances myocardial relaxation and reduces diastolic tone. However, the hypothesis that cGMP-mediated processes are the only mechanisms responsible for the NO-mediated contractile effects has subsequently been challenged. Several studies have demonstrated a dissociation between cGMP
concentrations and contractile state, because low concentrations of acetylcholine induced a negative inotropic effect even in the absence of any change in cGMP concentration. Recent reports show that under certain conditions, NO donors are able to enhance myocardial contractility, and that the basal intracellular production and release of NO significantly augments the Frank-Starling response in the isolated heart. It has been demonstrated that the augmentation of contractility by exogenous NO could be the result of an elevation of the intracellular levels of cAMP due to the cGMP-dependent inhibition of cAMP hydrolysis by cGMP-inhibited phosphodiesterase (cG-PDE or PDE type III). These observations, however, do not rule out the possibility of cAMP being increased by a direct (ie, GC/cGMP independent) modulation of the adenylcyclase (AC)/cAMP/cAMP-dependent protein kinase (PKA) pathway by NO.

Most reports on NO-mediated regulation of cardiac excitation and contraction have focused on the cGMP-dependent actions, whereas GC/cGMP-independent mechanisms have been largely unexplored. Relatively recently, a few publications have addressed “unconventional” pathways for NO signaling in the heart whereby NO may directly modulate protein function in a GC/cGMP-independent fashion, eg, via trans-nitrosylation of critical or regulatory thiols. Although NO certainly modulates cardiac contraction in part via cGMP/PKG-related mechanisms, considering the multiplicity of potential NO targets in the heart and the multifaceted effects of NO and NO-related compounds on cardiac contraction, it seems unlikely that NO exerts all of its physiologically relevant effects exclusively through activation of GC and PKG activity.

To examine whether NO potentially modulates the cAMP/PKA signaling pathway via GC-independent mechanisms, we studied the effects of the NO donor S-nitroso-N-acetylpenicillamine (SNAP), in the presence and absence of specific inhibitors of GC, PKG, and PKA, on contraction and the Ca2+ transient (Ca i) in isolated rat cardiac myocytes, in parallel with measurements of intracellular cGMP, cAMP, and AC activity. Parallel functional experiments were performed with 3-\([2\text{ - hydroxy } - 1\text{ - (1\text{-methylthyl}) } - 2\text{- nitrosohydrazinol} ] - 1\text{-propanamine (NOC5)}, an NO amine complex that serves as an NO donor chemically distinct from SNAP, and with NO scavengers, to establish that the biological effects were the results of the action of NO. The results from the present study suggest that NO is capable of evoking either positive or negative inotropic responses in cardiac myocytes depending on the concentration of exposure. Whereas the decrease in contractile response seen principally at higher levels of NO can be attributed mainly to a cGMP-dependent reduction in myofilament responsiveness to Ca2+, the enhanced contractile response at lower NO levels is due to increased intracellular cAMP levels that are mediated, at least in part, through a novel NO-dependent, GC/cGMP-independent activation of AC. Preliminary findings of the present study have been previously published in abstract form.

Materials and Methods

Cardiac Myocyte Isolation

Single cardiac myocytes were isolated via a previously described technique with minor modifications. Briefly, 2- to 4-month-old adult Sprague-Dawley rats (Charles River) were anesthetized with intraperitoneal sodium pentobarbital, and hearts were rapidly excised and perfused retrogradely with 25 mL nominally Ca2+-free bicarbonate buffer of the following composition (in mmol/L): NaCl 116.4, KCl 5.4, MgSO4 1.2, NaH2PO4 1.2, glucose 5.6, and NaHCO3 26.2, kept at 36±1°C and continuously gassed with 95% O2 and 5% CO2 to keep pH at 7.4. Perfusion was continued with a similar bicarbonate buffer containing 50 μmol/L Ca2+, 0.1% collagenase type B, 0.04 mg/mL protease XVI, and 0.1% BSA type V for ~20 minutes. The left ventricle was then mechanically dissociated, and myocytes were resuspended in a series of HEPES buffers with gradually increasing Ca2+ concentration. Cells were finally suspended in HEPES buffer containing (in mmol/L) NaCl 137, KCl 4.9, MgSO4 1.2, NaH2PO4 1.2, glucose 15, HEPES 20, and CaCl2 1.0, (pH 7.4) and stored at room temperature until use. Noncardiac myocytes comprised <1% of cells prepared in this manner. Cardiac myocyte viability was typically 70% to 80%. All experiments were replicated at least 3 times (or as specifically stated) and represent cells from at least 2 (or more) different myocyte preparations. Experiments were performed under a protocol approved by the National Institute on Aging Animal Care Committee.

Simultaneous Measurements of Contraction and Indo 1 Fluorescence

Changes in the Ca i, assessed by indo 1 fluorescence, and cell length were measured in isolated myocytes as previously described. In brief, for measurements of [Ca2+]i, myocytes were loaded with the membrane-permeable acetoxymethyl ester of indo 1 (indo 1-AM), using a 10-minute exposure to a 25 μmol/L solution in HEPES buffer at room temperature. After loading, cells were transferred to a Lucite chamber with a glass coverslip on the stage of an inverted microscope and were continuously superfused with HEPES buffer. Myocytes were chosen for study according to previously established criteria, ie, a rod-shaped appearance with clear striations and no membrane blebs, a negative staircase of twitch performance at rest, and the absence of spontaneous contractions. Cardiac myocyte contraction was produced via electrical field stimulation from rest, and the absence of spontaneous contractions. Cardiac myocyte contraction was produced via electrical field stimulation from rest, and the absence of spontaneous contractions. Cardiac myocyte contraction was produced via electrical field stimulation from rest, and the absence of spontaneous contractions. Cardiac myocyte contraction was produced via electrical field stimulation from rest, and the absence of spontaneous contractions. Cardiac myocyte contraction was produced via electrical field stimulation from rest, and the absence of spontaneous contractions. Cardiac myocyte contraction was produced via electrical field stimulation from rest, and the absence of spontaneous contractions. Cardiac myocyte contraction was produced via electrical field stimulation from rest, and the absence of spontaneous contractions. Cardiac myocyte contraction was produced via electrical field stimulation from rest, and the absence of spontaneous contractions. Cardiac myocyte contraction was produced via electrical field stimulation from rest, and the absence of spontaneous contractions. Cardiac myocyte contraction was produced via electrical field stimulation from rest, and the absence of spontaneous contractions. Cardiac myocyte contraction was produced via electrical field stimulation from rest, and the absence of spontaneous contractions. Cardiac myocyte contraction was produced via electrical field stimulation from rest, and the absence of spontaneous contractions. Cardiac myocyte contraction was produced via electrical field stimulation from rest, and the absence of spontaneous contractions.

Assessment of Myofilament Response to Ca2+

Changes in myofilament responsiveness to Ca2+ were assessed using the steady-state relation between cell length and [Ca2+]i in intact single cardiac myocytes tetanized by high-frequency (10 Hz) stimulation after exposure to thapsigargin (0.2 μmol/L for 15 minutes), as described previously. Thapsigargin discharges the sarcoplasmic reticulum and thus enables tetanization of otherwise intact myocytes. High-frequency electrical stimulation after thapsigargin treat-
ment results in the effective summation of the repetitive transmembrane flux of Ca\(^{2+}\) current (because of the absence of sarcoplasmic reticulum Ca\(^{2+}\) sequestration and periodic release) to achieve a steady-state level of myoplasmic Ca\(^{2+}\) substantially elevated above resting levels, at the point that Ca\(^{2+}\) influx is balanced by the rate of Ca\(^{2+}\) extrusion (via Na/Ca exchange). With this approach, the Ca\(_0\) can be reversibly clamped near peak systolic levels during the tetanic contracture for 10 to 60 seconds (or longer) and then rapidly returned to resting levels on cessation of electrical stimulation via normal Na/Ca exchange mechanisms. The steady-state levels of Ca\(^{2+}\) achieved during tetanization, moreover, can be systematically regulated by adjustments in the concentration of bathing Ca\(^{2+}\) and can be matched between protocols to effectively gauge changes in myofilament Ca\(^{2+}\) sensitivity. Thus, changes in the degree of cell shortening between tetani clamped at the same Ca\(_0\) level can be attributed to changes in the relative myofilament responsiveness to Ca\(^{2+}\).

**Determination of cGMP and cAMP**

Suspensions of freshly isolated adult rat cardiac myocytes were pretreated for 30 minutes at 23°C with either control or 1H-[1,2,4]oxadiazolo[4,3-a]quinoxalin-1-one (ODQ) 10 \(\mu\)mol/L containing buffers and subsequently challenged with the indicated concentration of SNAP for the next 20 minutes. The cells were then lysed with 1 mL ice-cold 0.6 mmol/L perchloric acid. The cell lysates (950 \(\mu\)L) were transferred to microcentrifuge tubes, and the pH was adjusted to 7.0 using K\(_2\)CO\(_3\). After centrifugation for 5 minutes at 8000 \(\times\) g, the supernatant was vacuum-dried and then recovered in 200 \(\mu\)L Tris/EDTA buffer. After addition of 0.15 mmol/L Na\(_2\)CO\(_3\) (20 \(\mu\)L) and 0.15 mmol/L ZnSO\(_4\) (20 \(\mu\)L) and incubation for 15 minutes on ice, the salt precipitate was removed by centrifugation for 5 minutes at 1200 \(\times\) g, and 50 \(\mu\)L of supernatant was assayed using either a cAMP [H] or cGMP [H] assay kit (Amersham). Cellular protein was measured using the Bradford method (Bio-Rad) with bovine \(\gamma\)-globulin as standard.

**Determination of AC Activity**

**Preparation of Purified Cardiac Sarcolemmal Membranes**

After mincing and washing with saline EGTA (154 mmol/L NaCl, 0.1 mmol/L EGTA), freshly excised cardiac ventricular tissue was suspended into 5 volumes of homogenizing buffer (HB) (in mmol/L: HEPES 10, MgCl\(_2\) 2, EGTA 0.1, and dithiothreitol 5; prepared the day of use). The suspension was homogenized 4 times for 10 seconds and centrifuged for 5 minutes at 1200 \(\times\) g for 15 minutes on ice, the salt precipitate was removed by centrifugation for 5 minutes at 1200 \(\times\) g, and 50 \(\mu\)L of supernatant was assayed using either a cAMP [H] or cGMP [H] assay kit (Amersham). Cellular protein was measured using the Bradford method (Bio-Rad) with bovine \(\gamma\)-globulin as standard.

**AC Enzyme Activity**

Two micrograms of membrane protein was added into a final volume of 200 \(\mu\)L containing (in mmol/L) HEPES 25, MgCl\(_2\) 0.4, GTP 5, dithiothreitol 0.15, NaCl 50, creatine phosphate 2.5, creatine phosphokinase 5 U, 3-isobutyl-1-methylxanthine (IBMX) 10, and alamethicin (1 \(\mu\)g/\(\mu\)L of membrane protein). The tubes were transferred to a 30°C water bath, and the reaction with a given dose of SNAP was started by adding 5 mmol/L ATP. After 15 minutes, the reaction was stopped by addition of 1.8 mL of 80°C to 90°C H\(_2\)O. Subsequently, cAMP was determined using the scintillation proximity assay according to the Amersham protocol furnished with the RPA 538 kit (Amersham).

**Determination of Free NO Concentration Produced by NO Donors SNAP and NOC5**

Free NO concentration was determined in the physiological buffer solutions containing either SNAP or NOC5 at the concentrations used in these experiments (90 minutes after preparation, 25°C) using a Sievers model 280 nitric oxide analyzer following the method recommended by the manufacturer (NO concentrations were also measured in the bathing solutions of some of the cardiac myocyte contraction experiments to ensure consistency of NO application). Calibration of the NO analyzer was obtained using reagent-grade NaNO\(_2\), prepared in nitrate-free, deionized water (standard solutions containing 10 mmol/L, 50 mmol/L, and 100 mmol/L and 1 \(\mu\)mol/L, 5 \(\mu\)mol/L, 10 \(\mu\)mol/L, and 100 \(\mu\)mol/L nitrate were used for calibration), in a nitrate-reducing reaction system of freshly prepared VCl\(_2\)/HCl at 90°C as prescribed by Sievers, which results in the quantitative conversion of nitrate (and nitrite) to NO. The resulting NO was measured by the detection of chemiluminescence from its reaction with ozone. Within-run and between-run reproducibility was \(\pm\)5%.

**Materials**

Collagenase type B was purchased from Boehhringer Mannheim Corp; isoproterenol from Winthrop Pharmaceuticals; protease X VI, N-acetylpenicillamine (NAP), and IBMX from Sigma Chemical Co; indo 1-AM from Molecular Probes Inc; BSA type V, bovine hemoglobin, SNAP, and NOC5 from Calbiochem Corp; Rp-8-CPT-cAMPS from Biolog; and ODQ from Biomol Research Laboratories Inc. All other chemicals were of the purest reagent grade available.

**Statistics**

All data are presented as mean \(\pm\)SEM. Comparisons within groups were made by paired and unpaired Student \(t\) test, and values of \(P<0.05\) were taken to indicate statistical significance.

**Results**

**Reversible Effects of SNAP on Cardiac Myocyte Contraction**

We assessed the effect of different concentrations of SNAP on contraction in electrically stimulated cardiac myocytes. Figure 1A shows representative examples of the opposing effects of high (100 \(\mu\)mol/L) and low (1 \(\mu\)mol/L) concentrations of SNAP on the unloaded contraction from 2 separate cardiac myocytes. At the high concentration (top tracing), SNAP induced a slowly evolving negative inotropic effect, which was maximal 15 to 20 minutes after administration of the drug, and this effect was completely reversible on washout. In contrast, low concentrations of SNAP induced a pronounced positive inotropic effect, which was entirely reversible on washout. Figure 1B shows the average data from these experiments, indicating the time course of the opposing contractile responses induced by high and low concentrations of SNAP. After 20 minutes of SNAP exposure, twitch amplitude was reduced by 24\% (n = 5) at the high concentration but was increased by 38\% (n = 5) at the low concentration. Thus, the dynamic range of contractile regulation induced by the different concentrations of NO used in these experiments is about two thirds of the basal contraction amplitude.

As control experiments, cells were exposed to the byproduct of SNAP after release of NO, NAP, at the same concentrations as used in the SNAP experiments. Figure 1C shows that neither NAP at 100 \(\mu\)mol/L nor NAP at 1 \(\mu\)mol/L had any effect on myocyte contraction, indicating that the various contractile effects seen with SNAP do not result from the thiol byproduct after the production of NO. Figure 1D demonstrates that the positive contractile effect seen in these experiments was indeed the result of NO, per se, insofar as that seen both with SNAP (1 \(\mu\)mol/L) and an unrelated NO
donor, NOC5 (1 μmol/L), was completely abolished by coincubation with the NO scavengers, oxyhemoglobin (10 μmol/L), and carboxy-PTIO (0.1 mmol/L; data not shown). The negative contractile effects seen at higher concentrations of both SNAP (100 μmol/L) and NOC5 (20 μmol/L) are similarly abolished by both oxyhemoglobin and carboxy-PTIO (data not shown). Similar results were obtained in at least triplicate observations in different cells under each of the conditions described above.

**Effect of SNAP on [Ca^{2+}]_i, cGMP, and cAMP**

Using indo 1–loaded cardiac myocytes, we investigated the effect of different concentrations of SNAP on contraction and Cai measured simultaneously. A representative example of the negative contractile effect of a high concentration of SNAP (100 μmol/L) and the associated Cai is depicted in Figure 2A. The progressive reduction in the twitch amplitude seen here was not associated with a decrease of the Cai, suggesting that SNAP (100 μmol/L) reduced the relative myofilament responsiveness to Ca^{2+}. Also shown in the figure are the results from parallel experiments comparing intracellular levels of cGMP and cAMP in the presence and absence of SNAP. SNAP (100 μmol/L) induced a large increase in cGMP (31±6% versus basal, P<0.05) but had no significant effect on cAMP levels. In contrast, a low concentration of SNAP (1 μmol/L) elicited a positive contractile response that was associated with a significant increase in the Cai amplitude (26±6% for n=5 cells, P<0.05; Figure 2B). This low concentration of SNAP induced only a modest increase in cGMP (12±3% versus basal, P<0.05) but produced a large increment in cAMP levels (35±9% versus basal, P<0.05). The average effects of high and low concentrations of SNAP on contraction and Cai amplitude and kinetics are provided in the Table.

**Effect of SNAP on Myofilament Responsiveness to Ca^{2+}**

The results shown in Figure 2 suggest that high concentrations of SNAP are associated with a diminished sensitivity of the myofilaments to Ca^{2+}. The effects of low concentrations of SNAP on myofilament sensitivity to Ca^{2+} are, however, more difficult to interpret, because both the amplitude of the Cai and the contraction change during the experimental protocol. To further characterize the effects of SNAP on myofilament responsiveness to Ca^{2+}, steady-state tetanic contractures (which enable the Cai to be reversibly maintained at reproducible levels near that of peak systole for periods of 10 to 60 seconds) were performed in the presence and absence of SNAP. Figure 3A shows a representative example of the effect of SNAP (100 μmol/L) on the steady-state myofilament response to Ca^{2+}. Despite the achievement of similar peak [Ca^{2+}]_i levels, in the presence of high concentrations of SNAP, steady-state cell contraction amplitude during the tetanus was significantly decreased, confirming that the phenomenon seen in Figure 2A was the result of a reduction in myofilament response to Ca^{2+}. In contrast, perfusion with a low concentration of SNAP (1 μmol/L) did not significantly affect either peak steady-state Ca^{2+} or shortening during the tetanus (Figure 3B), thus indicating that
low concentrations of SNAP do not exert an appreciable effect on myofilament responsiveness to \(\text{Ca}^{2+}\). Similar results were obtained in at least triplicate observations in different cells at each concentration of SNAP.

To assess whether the increase in cGMP induced by a high concentration of SNAP (ie, Figure 2A, right panel) activates PKG and is, in turn, responsible for the observed reduction in myofilament responsiveness to \(\text{Ca}^{2+}\), steady-state tetanic contractures were also performed in cardiac myocytes pretreated with the specific blocker of PKG KT 5823 1 μmol/L.

Figure 4 shows that inhibiting PKG activation with KT 5823 completely abolished the reduction in steady-state cell shortening. *\(P<0.05\) vs basal level.

### Effect of SNAP on Contraction and \(\text{Ca}^{2+}\) Transient Parameters of Single Rat Myocytes

<table>
<thead>
<tr>
<th></th>
<th>TA (% of Resting Cell Length)</th>
<th>(t_{1/2}) Contraction, ms</th>
<th>IFT Amplitude (410/490 nm)</th>
<th>(t_{1/2}) IFT, ms</th>
</tr>
</thead>
<tbody>
<tr>
<td>n</td>
<td>5</td>
<td>5</td>
<td>5</td>
<td>5</td>
</tr>
<tr>
<td>Control</td>
<td>5.7±0.3</td>
<td>354±14</td>
<td>0.168±0.01</td>
<td>260±15</td>
</tr>
<tr>
<td>SNAP 1 μmol/L</td>
<td>7.9±0.6*</td>
<td>350±14</td>
<td>0.211±0.02*</td>
<td>266±11</td>
</tr>
<tr>
<td>Control</td>
<td>6.1±0.4</td>
<td>340±10</td>
<td>0.155±0.03</td>
<td>264±18</td>
</tr>
<tr>
<td>SNAP 100 μmol/L</td>
<td>4.3±0.3*</td>
<td>311±12*</td>
<td>0.153±0.02*</td>
<td>256±15</td>
</tr>
</tbody>
</table>

*TA indicates twitch amplitude; \(t_{1/2}\) contraction, half-relaxation time of contraction; IFT amplitude, indo 1 fluorescent transient, an index of the \(\text{Ca}^{2+}\) transient amplitude; and \(t_{1/2}\) IFT, 50% relaxation of indo 1 fluorescent transient.

Values are mean±SEM. *Significant change vs control value (\(P<0.05\)).
ening during the tetany in the presence of high concentrations of SNAP (100 μmol/L). This protocol with KT 5823 has been shown to completely inhibit the negative contractile effect (of comparable magnitude to that seen here with SNAP 100 μmol/L) induced by 8-bromo-cGMP (50 μmol/L) in cardiac myocytes.21

**Effect of SNAP in the Presence of Inhibitors of GC**

To investigate whether cGMP-independent mechanisms participate in the positive inotropic effect elicited by a low concentration of SNAP, experiments were performed in the presence of the selective inhibitor of NO-sensitive GC36 ODQ (10 μmol/L). Figure 5A shows the effect of SNAP (1 μmol/L) on the contraction in a representative cardiac myocyte pretreated and continuously perfused with ODQ. In spite of the presence of ODQ, the positive inotropic response to SNAP (1 μmol/L) persisted (32±7% increase in twitch amplitude) together with an increased Ca i (20±6%) (n=6 cells). Perfusion with ODQ (10 μmol/L) alone did not significantly affect the baseline contraction or the Ca i. Figure 5C shows a representative example of the lack of effect of SNAP (1 μmol/L) in the presence of ODQ (10 μmol/L) on the steady-state myofilament response to Ca i (similar to the result in Figure 3B), which indicates that the positive contractile effect seen in Figure 5A is purely the result of the increase in magnitude of the Ca i (rather than a change in myofilament Ca i sensitivity).

The ability of ODQ to effectively inhibit cGMP production or to affect cAMP levels in response to SNAP was assessed in a parallel group of experiments in which cGMP and cAMP levels were measured in intact cardiac myocytes in the presence and absence of ODQ (10 μmol/L). That treatment with ODQ (10 μmol/L) was fully effective to block GC in these cardiac myocyte experiments is demonstrated by the fact it completely abolished the cGMP increase seen with 100 μmol/L SNAP: specifically, SNAP (100 μmol/L) alone increased basal cGMP (4.2±0.2 pmol/mg of protein) by 131±6%, whereas cGMP remained 98±4% of basal in the presence of SNAP (100 μmol/L) + ODQ (P<0.05). Figure 5B shows that ODQ completely abolished the increase in cGMP induced by SNAP 1 μmol/L, as expected, but did not affect the increased cAMP levels. Similar effects of ODQ on cGMP and cAMP levels were observed when higher concentrations of SNAP were used (data not shown). Moreover, preincubation of cardiac myocytes with ODQ was capable of trans-
forming an otherwise typical negative inotropic response of a high dose of SNAP (10 μmol/L) to a sustained positive inotropic response (Figure 6). These results indicate that low concentrations of NO are able to increase the Ca<sup>2+</sup> and contraction in intact cardiac myocytes in a cGMP-independent fashion. Although previous reports have suggested that the positive contractile response induced by NO donors could be the result of an elevation of the intracellular levels of cAMP (and PKA activation) due to cGMP-dependent inhibition of PDE type III (cGMP-inhibited phosphodiesterase or cGIPDE),<sup>8,9</sup> our results indicate the presence of an important, additional pathway whereby NO donors may increase cAMP and exert a substantial positive inotropic action, ie, through a GC/cGMP-independent mechanism.

Figure 6. Inhibition of GC with ODQ (10 μmol/L) to block the positive contractile response of SNAP (10 μmol/L) to a sustained positive response. The bar graph depicts the average change in contraction amplitude, expressed as a fraction of the basal contraction, after 20 minutes of exposure of cells to SNAP (10 μmol/L) either in the absence or presence of ODQ (10 μmol/L). SNAP alone significantly attenuated contraction amplitude by 26±4%, whereas in the presence of ODQ, SNAP induced a pronounced increase in contraction, 28±10% above baseline. *P<0.05 vs basal level.

Figure 5. Failure of ODQ 10 μmol/L to block the positive contractile response of SNAP 1 μmol/L. A, Typical continuous chart recording of contraction amplitude in response to ODQ (10 μmol/L) alone (top tracing) and SNAP (1 μmol/L) in the continued presence of ODQ (10 μmol/L) (bottom tracing). The superimposed tracings of cell length on the right show the lack of effect of ODQ (10 μmol/L) on baseline contraction (top) and the increase of contraction amplitude in response to a low concentration of SNAP (1 μmol/L) in the continued presence of ODQ (10 μmol/L) (bottom). B, Parallel measurements of cGMP and cAMP in response to SNAP (1 μmol/L) in the presence and absence of ODQ. The results are presented as fractions of the basal values of cGMP and cAMP, 4.2±0.2 and 6.2±0.4 pmol/mg of protein, respectively. ODQ (10 μmol/L) alone did not affect basal cGMP or cAMP levels (4.1±0.2 and 5.2±0.7 pmol/mg of protein, respectively). ODQ completely abolished the SNAP-induced increase in cGMP but had no effect on increased cAMP levels. *P<0.05 vs basal level. C, Representative example of the lack of effect of SNAP (1 μmol/L) in the presence of ODQ (10 μmol/L) on the steady-state myofilament response to Ca<sup>2+</sup> of a myocyte during tetanic contractions (10 Hz). This indicates that the positive contractile effect seen in panel A is purely the result of the increase in magnitude of the Ca<sub>i</sub>.

Effect of SNAP on AC Activity

Figure 8 illustrates the dose-response curve for the effect of SNAP on AC activity measured on cardiac sarcolemmal membranes. Basal AC activity was 92±6 pmol/mg of protein per minute. Low concentrations of SNAP (0.1 and 1 μmol/L, which produced 150 nmol/L and 500 nmol/L NO, respectively) showed significant increases of AC activity (19±6% and 16±4% above basal activity, respectively; P<0.05). At higher SNAP concentrations, AC activity was unchanged (at 10 μmol/L SNAP, which produced 1.4 μmol/L NO) or
preincubated with Rp-8-CPT-cAMPS 100 μM ODQ, Figure 9C and 9D, respectively). These results (increase in contraction in the presence of Rp-8-CPT-cAMPS for at least 1 hour before the experiment. In the continuous perfusion of intact rat cardiac myocytes continuously stimulated with SNAP, NAP, after NO release. Similarly, concentrations of NOC5 producing 0.5 μmol/L NO also enhanced myocyte contraction. Furthermore, these contractile effects were not induced by the thiol byproduct of NO donors, confirming these observations. Although the direction of the myocardial response clearly depends on the concentration of the NO donor used and on the level of intracellular cGMP reached, the principal novel finding of the present study is that a previously unrecognized GC/cGMP-independent mechanism involving the activation of AC is also critical in mediating this response.

Effect of SNAP in the Presence of Inhibitors of GC and PKA

To confirm that the persistent positive contractile response induced by low concentrations of SNAP in the absence of GC activation was mediated through cAMP-dependent PKA activation, experiments were performed using the cAMP analog Rp-8-CPT-cAMPS previously shown to effectively and specifically inhibit PKA-dependent processes. All cells were preincubated with Rp-8-CPT-cAMPS (0.1 μmol/L) and ODQ (10 μmol/L) for at least 1 hour before the experiment. In the continuous perfusion of the inhibitory cAMP analog Rp-8-CPT-cAMPS (100 μmol/L) and ODQ (10 μmol/L), low concentrations of SNAP were unable to elicit the positive inotropic response previously observed with SNAP alone. Figures 9A and 9B show representative examples of the failure of SNAP (1 μmol/L) to increase the contraction (or the associated Ca2+) of intact rat cardiac myocytes continuously perfused with Rp-8-CPT-cAMPS with and without ODQ, respectively (similar results were observed in 4 other preparations). Similarly, NOC5 (1 μmol/L) also fails to elicit an increase in contraction in the presence of Rp-8-CPT-cAMPS (±ODQ, Figure 9C and 9D, respectively). These results demonstrate that the positive inotropic effect of low concentrations of NO is, at least in part, independent of cGMP and is mediated by increased cAMP and PKA activation. In these same cells, as well as in parallel experiments, the ability of Rp-8-CPT-cAMPS to successfully inhibit PKA activation was routinely confirmed by its efficacy to inhibit the positive inotropic effect induced by isoproterenol. The mean increase in contraction amplitude induced by 1 μmol/L isoproterenol was 171±10% of the control value (n=4 cells), which was completely abolished by Rp-8-CPT-cAMPS (94±5% of control value; n=4 cells).

Discussion

The findings in the present study indicate that the exogenous administration of the NO donor SNAP can induce opposing effects on myocyte contraction, in agreement with recent reports. Additional experiments using NOC5, an NO donor chemically distinct from SNAP, confirm these observations. Although the direction of the myocardial response clearly depends on the concentration of the NO donor used and on the level of intracellular cGMP reached, the principal novel finding of the present study is that a previously unrecognized GC/cGMP-independent mechanism involving the activation of AC is also critical in mediating this response.

In rat cardiac myocytes, high concentrations of SNAP (producing ~14.7 μmol/L NO) induced large increases in cGMP and reduced myocyte contraction, whereas low concentrations of SNAP (producing in the range of ~0.1 to 0.5 μmol/L NO) induced only moderate increases in cGMP and enhanced myocyte contraction. Furthermore, these contractile effects were not induced by the thiol byproduct of SNAP, NAP, after NO release. Similarly, concentrations of NOC5 producing ~0.5 μmol/L NO also enhanced myocyte contraction. One can deduce that these contractile effects with
SNAP and NOC5 are authentic effects of NO, because they are completely reversible in the presence of the NO scavengers oxyhemoglobin and carboxy-PTIO. More importantly, our results show that whereas a cGMP/PKG-mediated diminishment of the contractile proteins to Ca\(^{2+}\) is largely involved in the negative inotropic response to high concentrations of SNAP, in isolated rat cardiac myocytes, is mediated primarily by a reduction in the myofilament sensitivity to Ca\(^{2+}\) rather than by a reduced Ca\(^{2+}\) availability at the myofilaments. In rat cardiac myocytes, administration of relatively high concentrations of the cGMP analog 8-bromo-cGMP (50 \(\mu\)mol/L) produced strikingly similar effects to those observed with high concentrations of SNAP (ie, a comparable negative inotropic response without any change of the Ca\(_i\)). Evidence for a cGMP-mediated reduction in myofilament responsiveness to Ca\(^{2+}\) has also been demonstrated in studies performed on skinned cardiac fibers. Previous reports also show that PKG activation induces a rightward shift of the tension/pCa relation in skinned single rat ventricular myocytes, compatible with a reduction in Ca\(^{2+}\) sensitivity of troponin C, and that PKG mediates phosphorylation of troponin I at the same site as that phosphorylated by protein kinase A, a mechanism known to reduce the affinity of troponin C for Ca\(^{2+}\). The precise underlying mechanism for the reduced myofilament responsiveness to Ca\(^{2+}\) induced by high concentrations of SNAP and increased cGMP was not directly addressed in our experiments. However, the ability of the PKG inhibitor KT 5823 to completely abolish the decrease in intact myocyte shortening observed during the tetanus in the presence of SNAP (100 \(\mu\)mol/L) (as it was also shown to do in this model in the presence of 8-bromo-cGMP) provides strong support for the concept that SNAP-induced PKG activation results in reduced myofilament responsiveness to Ca\(^{2+}\).

Although other reports have indicated that positive contractile responses are induced by low concentrations of NO donors and cGMP, the subcellular mechanisms involved have received relatively little attention. Some evidence suggests that the positive contractile response induced by NO donors could be the result of an elevation of the wide range of actions of NO (and of NO donors) on myocardial contractility, the subcellular mechanisms involved in the opposing contractile responses have not been completely elucidated.

In frog cardiac myocytes and in human atrial cells, cGMP has been shown to decrease Ca\(^{2+}\) influx through L-type channels by the stimulation of PDE type II (cAMP phosphodiesterase). In rat cardiac myocytes, however, cGMP has been shown to modulate L-type channel activity through a PKA-dependent pathway that does not involve changes in cAMP levels. Because Ca\(^{2+}\) entry through L-type Ca\(^{2+}\) channels is the principal trigger for Ca\(^{2+}\) release from the sarcoplasmic reticulum, the inactivation of L-type channels by NO/cGMP should have a significant effect on the amount of Ca\(^{2+}\) released to the myofilaments and therefore contribute to the negative inotropic response. This seems not to be an important mechanism in our model system, in which the administration of SNAP (100 \(\mu\)mol/L) produced a slowly evolving negative inotropic effect that was not associated with a reduction in the Ca\(_i\). Furthermore, in steady-state tetanic contraction experiments, SNAP (100 \(\mu\)mol/L) significantly decreased cell shortening during the tetanus, despite the achievement of similar peak [Ca\(^{2+}\)] levels compared with controls. Taken together, these results indicate that the negative inotropic response to high concentrations of SNAP, in isolated rat cardiac myocytes, is mediated primarily by a reduction in the myofilament sensitivity to Ca\(^{2+}\) rather than by a reduced Ca\(^{2+}\) availability at the myofilaments.

In rat cardiac myocytes, the absence of an inotropic response to SNAP is determined primarily by a cAMP/PKA-dependent, but GC/cGMP-independent, increase in the Ca\(_i\). Although recent reports have addressed the
intracellular levels of cAMP (and PKA activation) due to a cGMP-dependent inhibition of the PDE type III (ie, cGMP-inhibited PDE). Furthermore, low concentrations of cGMP (0.1 to 10 μmol/L) as well as of the NO donor SIN-1 (0.1 to 10 nmol/L) were found to have a stimulatory effect on L-type Ca²⁺ current. It seems likely, therefore, that the inhibition of cAMP degradation, mediated by the inactivation of the PDE type III after low cGMP levels accumulate, could result in the cAMP/PKA-dependent stimulation of transmembrane Ca²⁺ influx, increasing the trigger for the Ca²⁺-induced Ca²⁺ release process, and producing a larger Ca₅ and myocyte contraction. Nevertheless, in the current experiments, SNAP (1 μmol/L) was still capable of eliciting a pronounced positive contractile response (via increasing the Ca₅) in the presence of a background of tonically inhibited PDEs (via IBMX pretreatment; Figure 7). In this setting, it seems unlikely that cGMP accumulation, per se, could mediate additional PDE (type III) inhibition (and cAMP accumulation) beyond that tonically present, owing to the presence of IBMX, and this leads to the notion that the NO-mediated positive inotropy can occur, at least in part, via PDE-independent mechanisms.

However, a decrease in PDE activity is only one of the two possible major mechanisms capable of increasing cAMP in intact cells, the other being the “upstream” activation of AC. Our results show that the positive contractile response of SNAP (1 μmol/L) is indeed associated with a significant elevation of both basal cAMP and cGMP. Nevertheless, even when this basal cGMP increase was completely abolished by the presence of the selective inhibitor of soluble GC (ODQ) in the bathing solution (see Figure 5B), SNAP (1 μmol/L) was still able to induce a positive inotropic effect that was mediated by cAMP (Figures 5A and 9). In these settings, cAMP must be increased by a mechanism other than cGMP-mediated PDE inhibition, in all likelihood by a cGMP-independent activation of AC.

To evaluate the possible activation of AC by the NO donor SNAP, we determined AC activity in isolated sarcolemmal membranes from rat heart. SNAP induced a concentration-dependent biphasic response of AC activity, significantly increasing its activity at low doses (0.1 to 1 μmol/L) and either not changing (10 μmol/L) or marginally decreasing the activity at high doses (100 μmol/L). Consistent with our findings, recent studies provide evidence suggesting that NO released from endothelial cells can directly or indirectly activate AC in a cGMP-independent manner in isolated perfused rat kidneys. From the present experiments, we cannot ascertain whether NO activates AC directly or through a G protein (Gₛ/Gᵢ)–dependent pathway. Indeed, recent reports have demonstrated that NO can modulate G protein function. Lander et al. have demonstrated that treatment of intact human peripheral blood mononuclear cells with NO yielded membranes with enhanced GTPase activity, and that NO similarly enhanced the GTPase activity of pure, recombinant Gₛ, Gᵢ, and p2₁. Miyamoto et al. found in endothelial cells that NO can selectively inhibit G proteins of the Gₛ and Gᵢ family but not those of the Gₛ family, and that this modulation of G proteins could have a permissive action on the Gₛ/AC pathway. Thus, investigation of this question of the potential modulation of G protein signaling by NO in cardiac myocytes should prove directly applicable to the issue of how NO mediates the activation of AC and the positive inotropic effect.

The final remaining possibility to explain changes in AC activation in these experiments is that NO may modulate the activity of the β-adrenergic receptor (βAR) itself. It is noteworthy that the cAMP/PKA-mediated positive contractile effect of low concentrations of NO was associated neither with changes in the Ca₅ duration nor a lusitropic effect. Although this lack of effect on contraction and Ca₅, kinetics does not resemble the typical abbreviation of these parameters observed on β₁-adrenergic receptor (β₁AR) stimulation (which is also mediated by cAMP/PKA activation), on the other hand, it closely resembles the effects of another cAMP/ PKA signaling cascade activated by the specific stimulation of β₂-adrenergic receptors (β₂ARs). Like the NO-induced positive inotropic effect, β₂AR stimulation produces a slowly evolving increase in contraction amplitude (as compared with β₁AR stimulation) without affecting the kinetics of contraction or the Ca₅. The differential regulation of cardiac excitation-contraction coupling by β₁AR and β₂AR stimulation has been attributed to a unique functional compartmentalization of cAMP. Thus, it is tempting to speculate that a simple and unifying explanation for all the experimental results at low NO concentrations involves the specific activation of β₂ARs, or perhaps of G proteins specifically coupled to them, but this will require experimental verification.

The nature of the regulation of protein function by NO is an area of active investigation. Recent evidence suggests that thiols in proteins can recognize both nitrosative and oxidative events, which in turn may elicit distinct functional changes. For example, the extent of reversible poly-S-nitrosylation of thiol groups (R-S-NO) by NO was shown to correlate with the degree of activation of the cardiac Ca²⁺ release channel (ryanodine receptor), and this mode of post-translational sulphhydryl modification has been proposed as a general model of NO-mediated regulation of protein function. These mechanisms could plausibly operate at each of the effector proteins potentially involved in the NO-stimulated production of cAMP in these experiments, including the β-ARs, G proteins, and AC, each of which has multiple potential sites available for S-nitrosylation (ie, thiols of the many cysteine residues). Indeed, posttranslational modification of certain cysteines on these proteins has been shown to play an important role in several hormone signal transduction pathways, including the βAR system. For example, labile thioesterification via palmitoylation of a specific cysteine in the β₂ARs (and many other G protein–coupled receptors), as well as Gₛ, and the agonist-promoted dynamic turnover of the binding of this moiety, has been shown to influence the ability of these effectors to regulate the activation of AC. Thus, competition for these thiol sites by NO could exert significant regulatory effects on AC activation independently of, or in cooperation with, the usual agonist induction mechanisms.

Pinsky et al. showed that endogenous cardiac NO levels achieve dynamic, beat-to-beat oscillations on the order of...
micromolar magnitude, and that this phenomenon is amplified by ventricular loading, which could potentially serve an autoregulatory function. Indeed, it has recently been shown that intracardiac production and release of NO significantly augments the Frank-Starling response.26 These physiological NO levels in heart are thus comparable to that achieved in the present experiments using 0.1 to 1 μmol/L SNAP or 1 μmol/L NOCS (≈0.1 to 0.5 μmol/L NO) and associated with a positive inotropic effect. If the rate of denitrosylation of R-S-NO is sufficiently slow (ie, on the time scale of the cardiac cycle), then it is possible that these physiological NO oscillations achieve a functional integration or steady-state bound level, which in turn, could proportionately regulate the function of the effector protein. Such proportionate control of protein function via the degree of protein S-nitrosylation has been demonstrated for the cardiac ryanodine receptor, for example.29 It is thus possible that the activation of AC demonstrated in these experiments at levels of NO compatible with that achieved by the beating heart could indeed serve a physiological autoregulatory role. Notably, it has been shown that both myocardial cAMP55–58 and PKA 55 activities associated with that achieved by the beating heart could indeed serve an autoregulatory function. Indeed, it has recently been shown that intracardiac production and release of NO significantly mediated early relaxation of ventricular cardiac muscle. Circ Res. 1988; 62:1171–1174.


55. Wollenberger A, Babski EB, Krause EG, Genz S, Blohm D. Bogdanov EV. Cyclic changes in levels of cyclic AMP and cyclic GMP are the net result of the cardiac cycle. Biochem Biophys Res Commun. 1973;55:446–452.


Activation of Distinct cAMP-Dependent and cGMP-Dependent Pathways by Nitric Oxide in Cardiac Myocytes
Martin G. Vila-Petroff, Antoine Younes, Josephine Egan, Edward G. Lakatta and Steven J. Sollott

Circ Res. 1999;84:1020-1031
doi: 10.1161/01.RES.84.9.1020
Circulation Research is published by the American Heart Association, 7272 Greenville Avenue, Dallas, TX 75231
Copyright © 1999 American Heart Association, Inc. All rights reserved.
Print ISSN: 0009-7330. Online ISSN: 1524-4571

The online version of this article, along with updated information and services, is located on the World Wide Web at:
http://circres.ahajournals.org/content/84/9/1020

Permissions: Requests for permissions to reproduce figures, tables, or portions of articles originally published in Circulation Research can be obtained via RightsLink, a service of the Copyright Clearance Center, not the Editorial Office. Once the online version of the published article for which permission is being requested is located, click Request Permissions in the middle column of the Web page under Services. Further information about this process is available in the Permissions and Rights Question and Answer document.

Reprints: Information about reprints can be found online at:
http://www.lww.com/reprints

Subscriptions: Information about subscribing to Circulation Research is online at:
http://circres.ahajournals.org/subscriptions/