Identification of mechanisms involved in the relaxation of rabbit cavernous smooth muscle by a new nitric oxide donor ruthenium compound

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ABSTRACT

Purpose: The aim of this study was to evaluate the relaxation in vitro of cavernous smooth muscle induced by a new NO donor of the complex nitrosil-ruthenium, named trans-\([\text{Rut(NH}_{3})_{4}(\text{caffeine})(\text{NO})]\)C\(_{13}\) (Rut-Caf) and sodium nitroprusside (SNP).

Materials and Methods: The tissues, immersed in isolated bath systems, were pre-contracted with phenylephrine (PE) (1 µM) and then concentration-response curves (10\(^{-12}\) - 10\(^{-4}\) M) were obtained. To clarify the mechanism of action involved, it was added to the baths ODQ (10 µM, 30 µM), oxyhemoglobin (10 µM), L-cysteine (100 µM), hydroxicobalamine (100 µM), glibenclamide, iberotoxin and apamine. Tissue samples were frozen in liquid nitrogen to measure the amount of cGMP and cAMP produced.

Results: The substances provoked significant relaxation of the cavernous smooth muscle. Both Rut-Caf and SNP determined dose-dependent relaxation with similar potency (pEC\(_{50}\)) and maximum effect (Emax). The substances showed activity through activation of the soluble guanylyl cyclase (sGC), because the relaxations were inhibited by ODQ. Oxyhemoglobin significantly diminished the relaxation effect of the substances. L-cysteine failed to modify the relaxations caused by the agents. Hydroxicobalamine significantly diminished the relaxation effect of Rut-Caf. Glibenclamide significantly increased the efficacy of Rut-Caf (pEC\(_{50}\) 4.09 x 7.09). There were no alterations of potency or maximum effect of the substances with the addition of the other ion channel blockers. Rut-Caf induced production of significant amounts of cGMP and cAMP during the relaxation process.

Conclusions: In conclusion, Rut-Caf causes relaxation of smooth muscle of corpus cavernosum by means of activation of sGC with intracellular production of cGMP and cAMP; and also by release of NO in the intracellular environment. Rut-Caf releases the NO free radical and it does not act directly on the potassium ion channels.

INTRODUCTION

Human corpus cavernosum contains vascular smooth muscle, kept under tonic contraction induced by adrenergic excitation to maintain penile flaccidity (1). Nitric oxide (NO), the endothelium-derived relaxation factor discovered by Palmer and Moncada, is the main inhibitory neu-
rotransmitter that mediates penile erection in all animals (2). NO activates soluble guanylate cyclase (sGC), which induces production of cyclic guanosine monophosphate (cGMP) from guanosine triphosphate (GTP). The cGMP acts in intracellular effectors, like protein kinase G (PKG), which cause diminishment of intracellular calcium and disassociation of actin and myosin fibers, ultimately leading to relaxation of the smooth muscle (2).

NO donors are substances that release NO, either in vivo or in vitro. One of these donors, sodium nitroprusside (SNP), is a powerful vasodilator utilized in patients with hypertensive crisis. Nevertheless, it is extremely labile, induces tolerance and releases cyanide, which is toxic to the endothelium (3).

NO is a potent vasodilator synthesized by neurons, endothelial cells, leucocytes and platelets among others. It is released by the autonomous nerve terminals, and by the vascular and sinusoidal endothelium when stimulated by acetylcholine (4).

Synthesis of NO is catalyzed by NOS, which converts l-arginine and oxygen to l-citruline and NO. NOS exists as three isoforms in mammals: nNOS and eNOS are preferentially expressed in neurons and endothelial cells, respectively, and iNOS in virtually all cell types. All three NOS isoforms have been identified in the corpus cavernosum, with nNOS and eNOS being considered responsible for initiating and sustaining erection, respectively (5). Down-regulation of nNOS expression has been found in the corpus cavernosum of aging rats (6), a model in which corpus cavernous smooth muscle relaxation is impaired (7).

Endothelial dysfunction is present in a large group of patients with erectile dysfunction (ED) and co-morbidities like hypertension and diabetes. This syndrome is characterized by an endogenous NO production deficiency (8). About 56% of patients in this group show resistance to current ED treatment with phosphodiesterase 5 (PDE-5) inhibitors (9). Research for new drugs that increase bioavailability of endogenous NO is a permanent challenge.

Recently, new NO donor compounds with higher stability and less toxicity have been subject of research. One group of such substances, S-nitroso-glutathione (GSNO) and S-nitroso-N-acetylcysteine-etylester (SNACET), was utilized in studies with strips of human corpus cavernosum mounted in isolated bath systems, proving to be potentially useful for tissue relaxation (10).

SNP has already been utilized intracavernously in vivo, in both human and animals, to induce penile erection. In dogs and monkeys, SNP induced dose-related erection, without causing hypotension (11). In patients with ED, SNP provoked erection of shorter duration than did papaverine, without side effects - like hypotension, injection site pain, or priapism (11,12).

Ruthenium compounds, which are NO donors, have been tested in vitro and showed a similar relaxation to SNP on rat aorta smooth muscle (13). These compounds theoretically have the potential to be better than SNP because they do not release cyanide nor they react with the superoxide anion, forming peroxynitrite. However, further experimental studies are necessary to confirm this theory. Furthermore, they are stable at physiologic pH and soluble in water (13).

The aim of this study was to evaluate the relaxation in vitro of cavernous smooth muscle induced by a new NO donor of the complex nitrosyl-ruthenium, named trans-[Ru(NH3)4(caffeine)(NO)]C13 (Rut-Caf) and sodium nitroprusside (SNP).

MATERIALS AND METHODS

The study was approved by the local Committee of Ethics on Animal Use in Research. Adult male New Zealand rabbits, weight 2 - 3 kg, were used. After anesthesia, the penis was removed entirely, and placed in Krebs-Henseleit solution. Cavernous tissue was dissected free of connective tissue and albuginea, providing two corpus cavernosum strips (1 cm) from each penis.

These strips were mounted in isolated baths (10 mL) containing Krebs-Henseleit solution. This solution contains Na, K, Cl, Ca, MgSO4, HC03, PO4, glucose, albumin, and tromethamine (THAM) and was used to maintain tissues during experiments (14). The strips were aired with a mixture of O2 (95%) and CO2 (5%), pH 7.4, 37°C. Tissues were placed vertically, with tension of 1g. One extremity was connected to an isometric power...
transducer, while the other end was attached to a mobile unit that allowed tension adjustment.

The tissues rested for one hour. Tension was calibrated and the solution renewed every 15 minutes. Alterations on tension were registered on a polygraph (Gemini 7070, Ugo-Basile, Italy). Eight different experimental protocols were performed, as outlined next.

Experiment 1: After pre-contraction with phenylephrine (1 µM), rising concentrations (10^{-12} - 10^{-3} M) of SNP (as a control) and Rut-Caf were administered to the baths and relaxation concentration-response curves were obtained.

Experiment 2: To evaluate the NO liberation profile for Rut-Caf, oxyhemoglobin (10 µM) was added to the baths, 30 minutes before pre-contraction with phenylephrine (1 µM).

Experiment 3: To evaluate possible anion nitroxyln (NO-) liberation by Rut-Caf during the relaxation process, L-cysteine (100 µM), a specific NO- remover was added to the baths, 30 minutes before pre-contraction with phenylephrine (1 µM).

Experiment 4: To evaluate the contribution of metabolically activated potassium channels (K_{ATP}) on the relaxation process, one K_{ATP} blocker - glibenclamide (1 µM) - was added to the baths, 30 minutes before pre-contraction with phenylephrine (1 µM).

Experiment 5: To evaluate the contribution of high, medium and low conductivity calcium-activated potassium channels (K_{Ca}) on the relaxation, two K_{Ca} blockers - iberotoxin (1 µM) and apamine (0.1 µM) - were added to the baths, 30 minutes before pre-contraction with phenylephrine (1 µM).

Experiment 6: To evaluate the way NO is involved in the relaxation, hydroxocobalamin (0.1 mM), a NO remover, was added to the baths, 30 minutes before pre-contraction with phenylephrine (1 µM).

Experiment 7: To determine the activation of sGC by Rut-Caf, the agent 1H-[1,2,4]oxadiazele[4,3-d]quinoxalin-1-one (ODQ) - a sGC blocker - was added to the baths, 30 minutes before pre-contraction with phenylephrine (1 µM).

In all the experiments, relaxation concentration-response curves were obtained according to rising concentrations (10^{-12} - 10^{-3} M) of Rut-Caf.

Experiment 8: The RbCC strips were frozen in liquid nitrogen for dosage of cGMP and cAMP. The method of non-acetylation was utilized and samples were separated for dosage of proteins according to the Bradford method (1976) (15). Trichloroacetic acid (TCA) was added to the tissue macerate, resulting in a final TCA concentration of 10%. After centrifugation, the supernatant was washed with water saturated diethylic ether. This process was repeated six times. After the washings, the samples were dried in a nitrogen atmosphere at 60ºC, and then suspended again in the assay buffer of the immunoenzymatic kit for dosage of cGMP and cAMP.

The relaxant effect of the substances was measured from the maximal contraction plateau induced by phenylephrine, and expressed as percentages of contraction diminishment. The maximal effect (E_{max}) was considered as the maximal amplitude response induced by the relaxant agents on the concentration-response curves. The drug concentrations that induced 50% maximal relaxation (pEC_{50}) were determined after logarithmic transformation of concentration-response normal curves, and expressed as the negative logarithmic of values for each tissue (pEC_{50}). Percentages of contraction diminishment superior to 50% were considered significant.

The data were expressed as averages ± standard deviation. Statistical analysis was performed with ANOVA, followed by the Tukey-Kramer test. Values of p < 0.05 were considered significant.

RESULTS

The SNP E_{max} value was 100%, and its pEC_{50} was 6.9 ± 0.2. For Rut-Caf, E_{max} value was 72.6 ± 6.6%, and pEC_{50} was 6.8 ± 0.2. There was no significant difference in potency between the two substances (p = 0.851).

The relaxation curve induced by Rut-Caf was dislocated below with oxyhemoglobin, while the maximal response for Rut-Caf, in this setting, was 49.8 ± 6.5%.

Incubation of tissues with L-cysteine did not provoke any relaxation alterations. The values of pEC_{50} for Rut-Caf and Rut-Caf + L-cysteine were respectively 6.8 ± 0.2 and 6.9 ± 0.2 (p =
The $E_{\text{max}}$ value for Rut-Caf was 77% and for Rut-Caf + L-cysteine was 80% (Figure-1).

Glibenclamide modified potency and efficacy of Rut-Caf, as reflected by $pEC_{50}$ values for Rut-Caf of 5.2 ± 0.2 and Rut-Caf + glibenclamide of 7.3 ± 0.2 ($p = 0.016$). $E_{\text{max}}$ value increased from 68 ± 8.9% to 92 ± 7% ($p = 0.060$).

Tissue incubation with ibero-apamin did not interfere with Rut-Caf relaxations. The $pEC_{50}$ for Rut-Caf was 4.7 ± 0.2, compared to 4.8 ± 0.3 for Rut-Caf + ibero-apamin ($p = 0.779$) (Figure-2).

Hydroxocobalamin almost abolished the relaxant effects of Rut-Caf. The $pEC_{50}$ for Rut-Caf was 4.9 ± 0.1, while $pEC_{50}$ for Rut-Caf + hydroxocobalamin was 5.5 ± 0.2 ($p = 0.713$). As to $E_{\text{max}}$, values were respectively 85.3 ± 9.4% and 21.3 ± 2.2% ($p < 0.001$).

The relaxation induced by Rut-Caf was significantly inhibited by ODQ. This is evidenced by the $E_{\text{max}}$ values for Rut-Caf - 58 ± 9.2% - and Rut-Caf + ODQ - 34.6 ± 3.7% ($p = 0.038$) (Figure-3).

Both SNP and Rut-Caf produced cGMP in the cavernous tissue in amounts significantly higher than the basal control value of 8.26 pmol/mg (Emax = 80%) with potency similar to SNP; however, with lower maximum effect. The most probable hypothesis is that SNP acts on relaxation of vascular smooth muscle activating the sGC and as a hyperpolarizing agent through direct activation of potassium ion channels and Rut-Caf acts mainly on activation of sGC (16).

The soluble isoform sGC plays a pivotal role in erectile function because it provides the link between NO and cGMP, which represent the

**DISCUSSION**

This study evaluated a new ruthenium compound and its capability to promote relaxation of cavernous smooth muscle.

The substance Rut-Caf provoked significant relaxation of cavernous smooth muscle

![Figure 1 - Effect of Rut-Caf upon rabbit corpus cavernosum strips with and without pretreatment with 10 μM/L oxyhemoglobin, 1 mmol/L L-cysteine or 0.1 mmol/L hydroxocobalamin. Concentration/response curves were plotted for graded concentrations (10-12 to 10-3 M) of Rut-Caf. Results were expressed as average ± standard error based on seven experiments. The statistical significance was verified with ANOVA followed by Bonferroni’s test.](image-url)
extracellular and intracellular signaling molecules, respectively, in physiologic erection (17).

Bonaventura et al., studying substances that also belong to the nitrosyl-ruthenium complex, in rabbit aorta rings, demonstrated a maximal relaxation effect of 102% with pEC50 of 6.61 ± 0.09. These data confirm, as showed in this study, that ruthenium compounds are powerful vasodilators (13).

In vivo, NO arranges with hemoglobin creating a stable iron-nitrosyl complex that does not release NO. Bonaventura et al., evaluating the effects of ruthenium compounds in rabbit aorta rings in the presence of oxyhemoglobin, observed a decreased potency without significantly modifying its maximum effect, due to intracellular release of NO.

In this in vitro study, the addition of oxyhemoglobin to the organ baths reduced the maximum effect of Rut-Caf, but could not abolish it. Therefore, the relaxant effect of this substance is also probably due to the intracellular release of NO (13).

McDonald and Murad demonstrated that NO acts on the relaxation of vascular smooth muscle through activation of soluble guanylate cyclase, increasing the synthesis and bioavailability of intracellular cGMP (18). In this study, tissue incubation with a specific inhibitor of soluble guanylate cyclase (ODQ; 10 µM) abolished the re-
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This demonstrates the action of the substance studied on the NO-cGMP cell signaling system.

According to the studies with compounds of the group nitrosyl-ruthenium, it was observed that these substances act by means of donation of free NO and nitroxyl anion (NO⁻) (13). Utilizing a specific nitroxyl anion remover, L-cysteine, the potential of relaxation remained unchanged. However, the addition of the NO free radical remover, hydroxocobalamin, to the baths significantly decreased the relaxation induced by Rut-Caf. In contrast with Bonaventura et al. findings, Rut-Caf does not release the nitroxyl anion during relaxation. In consequence, probably its relaxation effect is related to release of the NO free radical.

Smooth muscle has neither a T-tubule system nor a well-developed sarcoplasmatic reticulum. Therefore extracellular calcium plays an

**Figure 3 - Concentration/response curves showing relaxation induced by Rut-Caf in rabbit corpus cavernosum strips with and without pre-treatment with 10 μM ODQ or 30 μM ODQ. Results were expressed as average ± standard error based on seven experiments.**

**Figure 4 - cGMP dosage of rabbit corpus cavernosum strips exposed to saline solution (control), 100 μM Rut-Caf or 3 μM SNP (positive control).**

**Figure 5 - cAMP dosage of rabbit corpus cavernosum strips exposed to saline solution (control), 100 μM Rut-Caf or 10 μM forskolin (positive control).**

*p < 0.05 vs. Rut-Caf with 10 μM ODQ pretreatment
*p < 0.01 vs. Rut-Caf with 30 μM ODQ pretreatment
important role, and calcium must enter the cytoplasm through the plasma membrane during an action potential.

A direct hyperpolarizing action of the Rut-Caf was not observed. The addition of calcium activated potassium ion channels of high, medium and low conductivity inhibitors did not modify the relaxation effect of Rut-Caf. When glibenclamide, which is an ATP dependent potassium channel blocker, was added to the baths, an increase on the efficacy of Rut-Caf was observed (pEC50 4.04 x 7.69, p < 0.05).

The results obtained in this study were similar to the findings of Lee and Kang in human corpus cavernosum (19). These researchers, studying the effect of relaxation of a NO donor - SIN-1, could observe an increase in the probability of opening of calcium activated potassium channels in an indirect way, through activation of soluble guanylate cyclase and activation of the cGMP protein kinase. However, it was not demonstrated direct action of SIN-1 on the ion channels studied.

The best explanation for the efficacy of Rut-Caf would be the action of glibenclamide as a reducing agent. Bates et al. showed that the bioactivation of NO in biological medium needs the presence of a reducing agent. In consequence, glibenclamide acting as a reducing agent increase the efficacy of Rut-Caf (3).

This hypothesis was supported by the spectroscopic profile. The product formed (UV-Vis) is compatible with the formation of the species Rut-Caf-H2O. Initially, there is a reduction of NO0 by glibenclamide with posterior liberation of the metal coordination sphere. In conclusion, this suggests that amplification in the presence of glibenclamide occurs due to reduction of NO+ present in the metal coordination sphere, increasing the concentration of NO0 in the reactive environment.

Although it is known that ion channels effectively act on the control of the basal tone of the cavernous smooth muscle (1), they do not participate in the relaxation process induced by Rut-Caf.

The production of intracellular cGMP from activation of sGC by NO in different tissues was initially demonstrated by Arnold et al. (20). The production of cGMP in corpus cavernosum of animals and humans by NO donors, during the relaxation process, was demonstrated in studies (10,21).

Similar to these studies, it was demonstrated that the production of cGMP induced by Rut-Caf in the relaxation of cavernous smooth muscle was significantly higher than that induced by saline solution and like that produced by SNP.

It was also observed production of cAMP induced by Rut-Caf during the relaxation process. Uckert et al. demonstrated the interaction of cGMP-cAMP cell signaling systems in the relaxation of cavernous smooth muscle (22). These authors demonstrated the presence of cAMP-specific phosphodiesterases, PDE-3 and PDE-5 in the cavernous endothelium. They also noted that the cAMP specific kinase PKA inhibited the relaxation induced by PDE-5 inhibitors, specific to cGMP, characterizing the integration of the cGMP-cAMP cell signaling systems on the cavernous smooth muscle relaxation. Although the mechanisms for that interaction are not clear, the authors have proposed that the increase in cAMP would be secondary to the drop in the level of cGMP, which would diminish the activity of PDE-3, increasing the level of cAMP. So, the regulation of cAMP-specific phosphodiesterases by cGMP could explain the increase of cAMP observed in this study, as a direct effect of Rut-Caf in the relaxation of cavernous smooth muscle.

Lindaman et al. studied the gallbladder smooth muscle in vitro. They noted that caffeine acting as an unspecific phosphodiesterases inhibitor 1 to 5, increased the production of cGMP and cAMP in the relaxation process (23). Similarly, the presence of caffeine in the structure of Rut-Caf could contribute to the increase of cAMP produced by that substance.

It should be emphasized that the results of this study were obtained in in vitro experimental model and future studies in vivo and clinical studies are necessary in order to confirm them.

CONCLUSIONS

In conclusion, Rut-Caf causes relaxation of smooth muscle of corpus cavernosum by means of activation of sGC with intracellular production of cGMP and cAMP; and also by release of NO
in the intra-cellular environment. Rut-Caf releases the NO free radical and it does not act directly on the potassium ion channels.

CONFLICT OF INTEREST
None declared.

REFERENCES