Restoring Nitric Oxide Cytosolic Calcium Regulation by Cyclic Guanosine Monophosphate Protein Kinase I Alpha Transfection in Coronary Endothelial Cells of Spontaneously Hypertensive Rats

Silvia Nistri\textsuperscript{a} Lorenzo Di Cesare Mannelli\textsuperscript{b} Luca Mazzetti\textsuperscript{b} Robert Feil\textsuperscript{c} Daniele Bani\textsuperscript{a} Paola Failli\textsuperscript{b}

Departments of \textsuperscript{a}Anatomy, Histology and Forensic Medicine and \textsuperscript{b}Pharmacology, University of Florence, Florence, Italy; \textsuperscript{c}Interfakultäres Institut für Biochemie Signaltransduktion – Transgene Modelle, University of Tübingen, Tübingen, Germany

Abstract

In microcoronary endothelial cells (RCEs) from spontaneously hypertensive rats (SHR), the nitric oxide (NO)/cyclic guanosine monophosphate (GMP)-dependent protein kinase I (cGKI) pathway cannot regulate the cytosolic calcium ([Ca\textsuperscript{2+}]_i) dynamic as in RCEs from Wistar Kyoto rats (WKY). We investigated the altered downstream NO target in SHR cells and, since cGKI expression was low, whether the re-expression of cGKI\textsubscript{α} in SHR RCEs could restore NO calcium responsiveness. We measured [Ca\textsuperscript{2+}]_i, dynamic by fura-2 imaging analysis and the cGKI level by RT-PCR and Western blot in SHR and WKY RCEs. Plasmids encoding for enhanced green fluorescence protein or cGKI\textsubscript{α}-enhanced green fluorescence protein were transiently transfected in SHR RCEs, and [Ca\textsuperscript{2+}]_i was evaluated. Angiotensin-II (AT-II) increased [Ca\textsuperscript{2+}]_i, in a concentration-dependent way in both strains. Whereas in WKY, endogenously produced NO and cyclic GMP analog decreased the AT-II-induced [Ca\textsuperscript{2+}]_i transient, they were ineffective in SHR RCEs. The cGKI level was low in SHR cells. However, after cGKI\textsubscript{α} re-expression, endogenous NO decreased the AT-II-induced [Ca\textsuperscript{2+}]_i transient, while endothelial NO synthase and cGKI inhibition prevented it. The low expression of cGKI in SHR accounts for the absent regulation of the agonist-induced [Ca\textsuperscript{2+}]_i transient by the NO/cyclic GMP pathway. Studies on cGKI in humans could contribute to a better understanding of cardiovascular pathologies.

Introduction

Endothelial cells accomplish a well-defined regulatory role in the maintenance of blood pressure homeostasis, platelet aggregation and vascular remodeling. Endothelial dysfunctions are involved in elevated blood pressure, atherosclerosis and thrombosis. Many endothelium-derived substances are involved in this delicate homeostat-

S.N., L.D.C.M. and L.M. contributed equally to this research.
cGKI and an insensitivity of the \([Ca^{2+}]_i\) spontaneously hypertensive rats (SHR), we reported a low thermore, in aortic smooth muscle cells isolated from expression of cGKI transient to NO/cyclic GMP/cGKI modulation. The re-activity being dependent on the decreased production of inositol 1,4,5-trisphosphate \([\text{GMP}_3]\)/cyclic GMP-dependent protein kinase type I (cGKI) is involved in this effect. The cGKI is a serine/threonine kinase able to phosphorylate many intracellular proteins, including calcium regulator proteins [4].

Indeed, early research in this field demonstrates that in Chinese hamster ovary cells transfected with a plasmid-encoding \(\alpha\) isomform of cGKI (cGKI\(\alpha\)), the \([Ca^{2+}]_i\) transient induced by thrombin is suppressed when the enzyme is activated by a cyclic GMP analog, this effect being dependent on the decreased production of inositol 1,4,5-trisphosphate [5]. In line with the effectiveness of cGKI in reducing the \([Ca^{2+}]_i\) transient induced by agonists, cGKI knock-out mice have altered the homeostatic control of blood pressure (hypertension) as well as increased platelet aggregation [6].

Other reports show that vascular smooth muscle cells isolated from cGKI knock-out mice cannot regulate the increase in \([Ca^{2+}]_i\) induced by agonists as cells from naive mice do [7]. Moreover, the re-expression of cGKI isoforms in smooth muscle cells of cGKI knockout mice restores NO regulation of \([Ca^{2+}]_i\) [7, 8]. Similar results have been obtained in Chinese hamster ovary cells, where stable transfection of the cGKI\(\alpha\) and, to a lesser extent, cGKI\(\beta\) can modulate the increase in \([Ca^{2+}]_i\) induced by thrombin, while in human coronary smooth muscle cells, the silencing of cGKI\(\alpha\) by RNA interference can potentiate \([Ca^{2+}]_i\), signaling induced by the same agonist [9]. Furthermore, in aortic smooth muscle cells isolated from spontaneously hypertensive rats (SHR), we reported a low expression of cGKI and an insensitivity of the \([Ca^{2+}]_i\) transient to NO/cyclic GMP/cGKI modulation. The re-expression of cGKI\(\alpha\) re-establishes NO/cyclic GMP/cGKI \([Ca^{2+}]_i\) regulation [10].

NO can also decrease \([Ca^{2+}]_i\) in endothelial cells and therefore regulates endothelium function [11]. In particular, the NO modulation of \([Ca^{2+}]_i\) can reduce the activation of eNOS and therefore its own production in an autocrine/paracrine mode. This negative feedback mechanism is believed to protect cells from the detrimental effect of excessive NO production [1, 11].

In microcoronary endothelial cells (RCEs) isolated from the heart of normotensive Wistar Kyoto (WKY) rats, NO (exogenously administered as NO donor) and atrial natriuretic peptide (that increases intracellular cyclic GMP by activating its guanylyl cyclase receptor) decreases the \([Ca^{2+}]_i\) transient induced by thrombin and bradykinin [12]. Also, the endogenously produced NO controls the \([Ca^{2+}]_i\) transient in WKY RCEs since eNOS inhibitors destroy this regulation.

On the other hand, in RCEs from SHR, \([Ca^{2+}]_i\) is insensitive to NO-mediated regulation. The lack of NO responsiveness in RCEs isolated from SHR is also observed when phosphodiesterases are inhibited and intracellular cyclic GMP is increased by atrial natriuretic factor [12]. Therefore, a downstream defective mechanism of this regulatory pathway could account for the alteration in \([Ca^{2+}]_i\), handling in SHR cells. On the other hand, cGKI activity and protein expression are reduced in whole heart homogenates [13] and cardiomyocytes [14] isolated from SHR. Therefore, we aimed to explore cGKI in more detail in RCEs and to test if its re-expression in SHR cells could restore NO \([Ca^{2+}]_i\) responsiveness. As AT-II plays an important role in hypertension, we decided to use this agonist to induce the \([Ca^{2+}]_i\) increase in RCEs, since the presence of AT-II receptors in RCEs and its effectiveness in inducing NO production has already been described [3].

Some of these data were presented at the Fourth International Conference on Relaxin and Related Peptide [15].

**Materials and Methods**

**Animals**

All animal manipulations were carried out according to the European Community guidelines for animal care (DL 116/92, application of the European Communities Council Directive 86/609/EEC). The ethical policy of the University of Florence conforms with the Guide for the Care and Use of Laboratory Animals of the US National Institutes of Health (NIH Publication No. 85-23, revised 1996; University of Florence assurance number: A5278-01). Formal approval to conduct the experiments described was obtained from the Animal Subjects Review Board of the University of Florence. For all experiments described herein, 12- to 14-week-old male normotensive WKY (310.5 ± 6.95 g) and age-matched SHR (311.7 ± 9.26 g) were used (Charles River, Italy). Four rats were housed per cage (size 26 × 41 cm); animals were fed with standard laboratory diet and tap water ad libitum and kept at 23 ± 1°C with a 12-hour light/dark cycle, light at
7 a.m. Animals were anesthetized before sacrifice by cervical dislocation and hearts were isolated as described [12]. No differences between WKY and SHR heart/body ratios were measured, as previously described [16].

**Isolation and Culture of Rat Coronary Endothelial Cells**

Hearts were digested with 0.1% collagenase in Krebs–Henseleit solution; the obtained cell suspension was treated with trypsin (0.05% in phosphate-buffered saline for 30 min at 37°C) and plated in culture flasks for 4 h in complete RCE medium (M-199 containing 10% fetal bovine serum and 10% newborn calf serum, 250 IU ml⁻¹ penicillin G and 250 μg ml⁻¹ streptomycin). Then, non-adherent cells were removed by washing and adherent endothelial cells were grown until confluence. The obtained cells were characterized as previously described [12, 17] by their positive uptake of acetylated low-density lipoproteins (1,1'-dioctadecyl-3,3',3'-tetramethyl-indocarbocyanine perchlorate complex; Biochemical Technologies Inc., Stoughton, Mass., USA). For all experiments, cells were used at the first passage in culture. Twenty-eight different cell lines (10 from WKY and 18 from SHR) were used for these experiments.

**Cytosolic Intracellular Calcium Determination**

\[ \text{[Ca}^{2+}]_i \text{] was measured using a ratiometric image analysis system} \]

For \([Ca^{2+}]_i \) determination, two excitation wavelengths (340 and 380 nm) and a changeable dichroic filter block. For \([Ca^{2+}]_i \) determination, two excitation wavelengths (340 and 380 nm), a 400-nm dichroic mirror (DM 400) and a 510 nm barrier filter (BA 510) were used. Video images were obtained with an extended ISIS-M camera (Photonic Science, Robertsbridge, UK). Ten to 15 cells were analyzed in each microscopic field. Following extended ISIS-M camera (Photonic Science, Robertsbridge, UK). For all experiments, cells were used at the first passage in culture. Twenty-eight different cell lines (10 from WKY and 18 from SHR) were used for these experiments.

**Reverse Transcription PCR for cGKI mRNA**

The mRNA level for cGKI was assayed by reverse transcription PCR (RT-PCR) in WKY and SHR RCEs and in SHR cells transfected with cytomegalovirus immediate early promoter fused with enhanced green fluorescent protein plasmid (pcMV-EGFP) or pcMV-cGKI-EGFP (whole-cell culture extract; see below for plasmid transfection). 500 ng of total RNA was reverse transcribed and amplified with SuperScript™One-Step™ RT-PCR System (Invitrogen Life Technologies), as previously described [10]. The following rat gene specific primers were used: cGKI mRNA (NM_001055731), forward 5’-CATACATTGGGACACGAGACA-3’ and reverse 5’-GCAGACGGTCTTAACACACA-3’; β-actin mRNA (NM_031144), forward 5’-TTACACCTGAAAAGATGACC-3’ and reverse 5’-AGAGGTCTTTACGGATG-TCA-3’. The expected length of the amplified fragments was 233 and 539 bp for cGKI and β-actin, respectively. Densitometric analysis was conducted by measuring the bands as well as the smear of each lane by means of Scion Image analysis software. β-Actin normalization was performed for each result.

**Western Blot for Soluble Guanylyl Cyclase and cGKI**

Confluent RCEs were lysed in cold buffer (Tris/HCl, pH 7.4, 10 mM, NaCl 10 mM, MgCl₂ 1.5 mM, NAD, EDTA 2 mM, phenylmethylsulfonyl fluoride, 1 mM, Triton X-100 1%, leupeptin 20 μg/ml, pepstatin 1 μg/ml, Pefabloc® 1 mg/ml, aprotinin 2.5 μg/ml) and centrifuged. Freshly isolated ventricles from SHR and WKY hearts were immediately frozen in liquid nitrogen, homogenized in cold lysis buffer, and the obtained suspension was centrifuged. Seventy-five micrograms of supernatant proteins were electrophoresed by SDS-PAGE (7.6% polyacrylamide), blotted onto nitrocellulose membranes (Amersham, Cologno Monzese, Italy) and peroxidase-labeled goat anti-rabbit antibodies (1:10,000; Vector, Burlingame, Calif., USA) and peroxidase-labeled rabbit anti-chicken antibodies (1:1,500,000; Enzo Life Sciences, Lausen, Switzerland) and peroxidase-labeled rabbit anti-chicken antibodies (1:1,500,000; Enzo Life Sciences, Lausen, Switzerland) and peroxidase-labeled goat anti-rabbit antibodies (1:10,000; Vector, Burlingame, Calif., USA). For soluble guanylyl cyclase immunoblot, chicken polyclonal anti-soluble guanylyl cyclase (α₁-β₁ heterodimer) antibodies (1:500; Enzo Life Sciences, Lausen, Switzerland) and peroxidase-labeled rabbit anti-chicken antibodies (1:1,500,000) were used. Membranes were also immunostained with rabbit polyclonal anti-β-actin antibodies (1:20,000). Immune reaction was revealed using enhanced chemiluminescence (Amersham). Densitometric analysis was conducted by measuring the bands of each lane by means of the Scion Image analysis software. β-Actin normalization was performed for each result.

**Transient Transfection of cGKIα Isoform Plasmid**

The pcMV-cGKIα-EGFP vector was engineered to obtain a construct encoding cGKIα as a fusion to the N-terminus of EGFP [7]. pcCMV-EGFP encoded EGFP (Clontech-Celbio, Pero Milanese, Italy) only. These mammalian expression plasmids confer kanamycin resistance and contain a CMV immediate early promoter. pcMV-cGKIα-EGFP and pcMV-EGFP were transformed in a DH5α competent Escherichia coli strain (Novagen-Merck, Darmstadt, Germany). A single colony from a plate was used to inoculate 250 ml of Luria-Bertani medium containing 30 μg/ml kanamycin, and the culture was incubated for 16 h at 37°C. Bacteria were collected by centrifugation at 4°C for 15 min at 6,000 g. Amplified plasmid DNA was purified by QIAfilter Plasmid MaxiPrep (Qiagen, Milan, Italy). The fusion protein possesses enzymatic activity similar to the native form [7].

The described expression plasmids encoding cGKIα-EGFP protein or EGFP alone were transfected in semiconfluent RCEs from SHR using Lipofectamine™ (Invitrogen Life Technologies) in a ratio of 6 μg DNA:151 μg lipofectamine, and cells were grown for an additional 72 h. Cells were loaded with fura-2, and [Ca2+]i was measured as described in the previous paragraph. In each optical field examined for [Ca2+]i measurement, we identified cells expressing plasmids (positive cells) by looking at their EGFP fluorescence using an excitation wavelength of 490 nm, emission of 510 nm and a dichroic mirror at 500 nm. No interferences between fura-2 and EGFP fluorescence were detected.

Unless otherwise specified, all reagents were purchased from Sigma-Aldrich (Milan, Italy) and were of the highest analytical grade.
Data are reported as means ± SEM; n indicates the number of different cell lines analyzed. Statistical analysis was performed using one-way ANOVA followed by the Student-Newman-Keuls test. A p value ≤0.05 was considered as significant.

Results

Effect of AT-II in WKY and SHR RCEs
In basal conditions, [Ca^{2+}]_{i} was 119.5 ± 5.76 (n = 10) and 137.1 ± 7.42 nM (n = 16) in WKY and SHR cells, respectively. As previously described in RCEs isolated from Wistar rats [19], AT-II concentration-dependently induced a transient increase in [Ca^{2+}]_{i} in RCEs from WKY and SHR (fig. 1). In the absence of extracellular calcium, this [Ca^{2+}]_{i} transient was reduced, while after incubation with thapsigargin, the [Ca^{2+}]_{i} signal was practically absent (fig. 2a). The effect of AT-II was prevented by incubating cells with 10^{-6} to 10^{-8} M irbesartan (fig. 2b), suggesting that AT-II increased [Ca^{2+}]_{i} in RCEs by activating its type 1 receptor. A similar pathway of [Ca^{2+}]_{i} signaling was found in SHR cells (data not shown).

By comparing the AT-II-induced [Ca^{2+}]_{i} signaling in WKY and SHR cells, we revealed that the [Ca^{2+}]_{i} transient was characterized by a slightly higher peak in RCEs from SHR (fig. 3a, b) followed by a much slower decline to baseline, as demonstrated by the significant enhancement of the decay time as compared to WKY cells (fig. 3a, c). In RCEs from WKY, the rapid decline in the AT-II-induced [Ca^{2+}]_{i} transient was significantly slowed by incubation with the specific NOS inhibitor L-NAME (N\(^{\text{w}}\)/H\(_{9275}\)-nitro-L-arginine methyl ester HCl; 10^{-4} M, 10 min, 37 °C; fig. 3c). Of note, NOS inhibition did not change the [Ca^{2+}]_{i} transient in RCEs from SHR (fig. 3c), and in this experimental condition, no significant differences between WKY and SHR cells were observed. Similar behavior was also observed when WKY cells were pretreated with the cGKI inhibitor KT-5823 (10^{-6} M, 10 min, 37 °C) before administration of AT-II. KT-5823 increased the decay time in WKY, but did not modify it in SHR cells. Conversely, pre-treatment with the membrane-permeable cyclic GMP analog DBC (N\(^{2},N^{\prime}\)/H\(_{11541}\)-O-dibutyrylguanosine 3',5'-cyclic monophosphate sodium salt; 10^{-6} M, 10 min, 37°C) significantly reduced the [Ca^{2+}]_{i} transient peak and decay time in WKY; being ineffective in SHR cells to a lesser extent, the NO donor S-nitroso-N-acetylpenicillamine (SNAP) slightly reduced both the [Ca^{2+}]_{i} transient peak and decay time in WKY (not significant) and was completely ineffective in SHR cells (fig. 3b, c). Since the influ-

![Fig. 1.](image1) ![Fig. 2.](image2)

*Fig. 1.* Concentration-dependent effect of AT-II on [Ca^{2+}]_{i} in RCEs isolated from WKY and SHR. Typical time course of AT-II (10^{-6} to 10^{-8} M) induced calcium transient in WKY and SHR RCEs. AT-II was administered at time 0. Traces are the mean of at least 10 cell recordings. [Ca^{2+}]_{i} was measured using fura-2 fluorescence applied to a ratiometric image analysis system. For more details, see Materials and Methods.

*Fig. 2.* Characterization of AT-II signal in rat RCEs. Typical time course of 10^{-6} M AT-II signal in WKY RCEs. AT-II was administered at time 0. **a** AT-II was administered in control conditions, extracellular calcium-free medium ([Ca^{2+}]_{\text{out}} = 0) or in the presence of 30 nM thapsigargin (administered at the arrow). **b** AT-II was administered in control conditions or in the presence of 10^{-6} to 10^{-8} M irbesartan. Irbesartan was preincubated for 10 min before AT-II. Traces are the mean of at least 10 cell recordings. [Ca^{2+}]_{i} was measured using fura-2 fluorescence applied to a ratiometric image analysis system. For more details, see Materials and Methods.
ence of self-produced endogenous NO could potentiate the effectiveness of DBC, we performed experiments in which DBC was coincubated with the eNOS inhibitor L-NAME. When WKY cells were contemporaneously incubated with L-NAME and DBC, both [Ca\(^{2+}\)] transient peak and decay time were decreased as compared to L-NAME alone. All these results showed that NO/cyclic GMP/cGKI pathway modified the AT-II-induced [Ca\(^{2+}\)] transient in WKY, whereas it was ineffective in SHR RCEs (fig. 3b, c).

**cGKI Level in RCEs**

As shown by RT-PCR, the mRNA for cGKI was lower in SHR than in WKY (fig. 4a). Densitometric analysis showed that the mRNA level was significantly reduced in SHR RCEs (fig. 4b). Similar results were obtained by monitoring cGKI protein by Western blot analysis (fig. 5a). As shown by the densitometric analysis (fig. 5b), the enzyme was significantly less expressed in SHR as compared to WKY cells. Also, in freshly isolated heart ventricles, cGKI expression was significantly lower in SHR than in WKY (fig. 5). On the other hand, soluble guanylyl cyclase in RCEs of WKY and SHR was not differently expressed as measured by densitometric analysis of Western blot (4.3 ± 0.45 and 4.8 ± 0.37 arbitrary units in WKY and SHR, respectively; n = 4).
Fig. 4. mRNA level of cGKI in WKY and SHR RCEs. a mRNA level for cGKI was assayed by RT-PCR in both naïve (SHR), cGKIα-transfected SHR (SHR cGKIα-T) and naïve WKY RCEs. The first gel lane shows molecular weight markers. b Densitometric analysis (means ± SEM) of the mRNA level of 4 separate RCE experiments. β-Actin normalization was performed for each result. a p < 0.05 versus other groups (one-way ANOVA followed by Student-Newman-Keuls test); b p < 0.05 versus SHR cGKIα-T.

Effect of cGKIα-EGFP Transfection on [Ca2+]i

Dynamics in SHR

Consistently with these results, we transfected SHR RCEs with a plasmid encoding for cGKIα-EGFP or EGFP alone. In the whole cell culture extract, the transfection of cGKIα-EGFP significantly increased cGKI mRNA quantity, carrying it to a higher level than in WKY (fig. 4). As identified by their EGFP fluorescence, positive cells were 35–40% of the total cell number. A representative image of transfected cells is depicted in figure 6.

Thereafter, [Ca2+]i was studied in cGKIα-EGFP (cGKIα-positive) and EGFP-positive cells. The re-expression of cGKIα consistently modified the [Ca2+]i transient induced by AT-II (fig. 7). In fact, in cGKIα-positive cells, the Δ[Ca2+]i induced by AT-II was significantly decreased as compared to EGFP-positive (fig. 7a) and -naïve (fig. 3b) RCEs. Similarly, the decay time was significantly reduced in cGKIα-positive RCEs (fig. 7b). Conversely, EGFP-positive cells (fig. 7) behaved as naïve SHR RCEs. In order to test whether the decrease in the AT-II-induced [Ca2+]i transient in cGKIα-positive cells was dependent on the endogenous production of NO, we performed experiments in cells preincubated with the NOS inhibitor L-NAME (10^-4 M, 10 min, 37°C). As shown in figure 7, in cGKIα-positive cells, preincubation with L-NAME significantly increased Δ[Ca2+]i (fig. 7a) and decay time (fig. 7b) as compared to their cGKIα-positive control cells, whereas both parameters remained unchanged in EGFP-positive RCEs. Similarly, preincubation with KT-5823 modified the [Ca2+]i transient in cGKIα-positive RCEs but was ineffective in EGFP-positive cells. Moreover, although DBC did not significantly modify the [Ca2+]i transient in cGKIα-positive cells, it counteracted the effect of L-NAME incubation. Also, SNAP can slightly decrease the [Ca2+]i transient and decay time in cGKIα-positive cells.

Discussion

Our data demonstrate that the inadequate expression of cGKI accounts for the absent regulation of [Ca2+]i by the NO/cyclic GMP/cGKI pathway in SHR RCEs. Moreover, the re-expression of cGKIα is sufficient to restore NO/cyclic GMP/cGKI control of the AT-II-induced [Ca2+]i transient, thus underlying the central role of cGKI in calcium signaling.

cGKI has been less extensively studied in endothelium than in other tissues although several authors report its expression and function in primary cultures of microvascular endothelial cells [20, 21]. Among other functions, cGKI can reduce NO production in endothelial cells by means of a negative feedback mechanism [11]. Indeed, NO (produced in endothelial cells by the calcium-dependent enzyme eNOS) decreases agonist-induced [Ca2+]i transients through cGKI activation. Therefore, by reducing [Ca2+]i, NO can modulate its own production by a negative autocrine loop [11]. This autoregulation represents an important protective mechanism from the detrimental effects of excessive [Ca2+]i and NO. Specifically, in aortic endothelial cells, AT-II can stimulate the production of NO and peroxynitrite [2]. It is well established that peroxynitrite can interact with lipids, DNA, and proteins through direct oxidative reactions or indirect, radical-mediated mechanisms and its excessive production can induce cell necrosis or apoptosis [22, 23].

Several authors have already described AT-II activity on endothelial cells [2, 3, 19]. As reported by Bayraktutan...
and Ulker [3], AT-II increases NO production in rat RCEs by acting through AT1 receptors as its action is prevented by valsartan. Therefore, due to its relevant role in hypertension, AT-II appears as an adequate agonist for the study of the role of the NO/cyclic GMP/cGKI pathway in SHR RCEs.

Our research shows that AT-II induces a concentration-dependent increase in $[\text{Ca}^{2+}]_i$ in both WKY and SHR RCEs. This $[\text{Ca}^{2+}]_i$ signal is mainly dependent on the discharge of intracellular calcium stores followed by an influx from extracellular milieu. Indeed, incubation with thapsigargin strongly reduces it, while in the absence of extracellular calcium, the $[\text{Ca}^{2+}]_i$ transient is reduced but still maintained. This $[\text{Ca}^{2+}]_i$ signal characterization is in line with that already described by Nilius and Droogmans [1]. AT-II acts primarily on AT1 receptors, since it is antagonized by irbesartan in both strains. Also, the NO production as described by Bayraktutan and Ulker [3] is mainly antagonized by valsartan, demonstrating a large involvement of AT1 in this effect.

In WKY cells, AT-II-induced $[\text{Ca}^{2+}]_i$ increase is strictly regulated by the NO/cyclic GMP/cGKI pathway, since NOS and cGKI inhibition increases the $[\text{Ca}^{2+}]_i$ transient (mainly the decay time), while DBC and to a lesser degree SNAP decreases it. According to Kasseckert et al. [24], cGKI$\alpha$ activation can increase $[\text{Ca}^{2+}]_i$ sequestration in the endoplasmic reticulum in endothelial cells isolated from rat coronary circulation. This effect is dependent on

![Fig. 5. cGKI protein expression in WKY and SHR RCEs and freshly isolated heart ventricles. a] Protein expression for cGKI in confluent RCEs and freshly isolated ventricles. b] Densitometric analysis (means ± SEM) of cGKI protein levels of 4 different experiments. β-Actin normalization was performed for each result. $^a$ p < 0.05 (one-way ANOVA), SHR versus WKY.

![Fig. 6. Image of SHR RCEs transfected with cGKIα-EGFP. The fluorescence of EGFP was monitored at 490 nm excitation and 510 nm emission, as described. Bar = 50 μm.]
the phosphorylation of phospholamban which, in turn, can increase endoplasmic reticulum calcium ATPase pump turnover. This mechanism may explain the high responsiveness of decay time to the NO/cyclic GMP/cGKI pathway observed in WKY RCEs. On the contrary, in SHR RCEs (where cGKI expression is low), the NO/cyclic GMP/cGKI pathway does not modulate $[\text{Ca}^{2+}]_i$ transient. The low expression of cGKI seems to be a specific feature in the NO/cyclic GMP/cGKI pathway, since eNOS is not reduced in these cells [12], nor is soluble guanylyl cyclase (present data).

In order to test if the low expression of cGKI can account for this difference in calcium parameters, we have transfected cGKI$\alpha$ in SHR RCEs. The re-expression of cGKI$\alpha$ in SHR cells fully restores NO/cyclic GMP/cGKI regulation of the $[\text{Ca}^{2+}]_i$ transient, thus demonstrating that cGKI is the missing step in SHR signaling. In smooth muscle cells isolated from SHR aortas, our research group has already described a similar impairment of NO/cGMP/cGKI-mediated calcium handling [10]. It should be noted that, according to the present data, this alteration is not restricted to a single cell type or organ, but may represent a generalized feature.

The consequence of NO/cyclic GMP/cGKI pathway alteration in SHR RCEs could be at least as important as in vascular smooth muscle cells. Indeed, if the inactivity of NO-dependent calcium regulation in SHR vasculature is mainly related to contraction and hypertension, it may imply an over-functionality of eNOS by a reduced inhibitory feedback pathway in RCEs. This pathological NO production leads to toxicity on RCEs (autocrine loop) and cardiac myocytes (paracrine loop) [23]. The cGKI might influence eNOS activity not only by regulating $[\text{Ca}^{2+}]_i$, but also by direct phosphorylation of eNOS [25]. However, the precise role of cGKI in eNOS activation is still unclear and deserves further investigation [26].

Other pathways can regulate intracellular $[\text{Ca}^{2+}]_i$ signaling in many cell types. In particular, a complex inter-
action between cyclic AMP and cyclic GMP through their respective kinases deserves deep investigation. The precise understanding of how these cross-talk mechanisms are regulated and coordinated in endothelial cells is an important direction for future research.

**Conclusions**

According to these data, we can hypothesize that in SHR RCEs, a high quantity of biologically inactive NO is produced, which reacts with other oxygen-reactive species to form peroxynitrite. Peroxynitrite is implicated in heart and vascular dysfunctions through multiple mechanisms including lipid peroxidation, inhibition of mitochondrial respiration, reduction in cardiomyocyte contraction and apoptosis [23]. Therefore, the low expression of cGKI in SHR RCEs may be responsible for cardiac pathologies other than blood pressure regulation.

The role of cGKI in hypertension has been extensively investigated in hypertension and related diseases. Target disruption of the cGKI gene in mice induces a hypertensive phenotype [27]. A reduced mRNA expression of cGKI has been described in aortic rings of 6-week-old SHR [28]. Since at this age SHR are not hypertensive, it could be concluded that a low cGKI expression can contribute to the genetic predisposition of SHR to hypertension. Moreover, both in SHR and in deoxycorticosterone acetate (salt) rats, a mineralocorticoid hypertensive rat model, the increase in aortic cGKI expression induced by hemin is associated with a significant decrease in blood pressure, further suggesting a strict relationship between cGKI and blood pressure [29]. Similarly, in stroke-prone SHR, the AT-II type 1 receptor antagonist valsartan reduces blood pressure and increases the cGKI expression in mesenteric arteries [30]. However, it remains to be established if the decrease in blood pressure in these hypertensive animal models is the cause or the effect of cGKI protein increase. A different feature has been observed in rats infused for 7 days with AT-II (1 mg/kg/day): blood pressure is dramatically increased, while in aortas, cGKI protein expression remains unchanged [31].

Therefore, the relationship between cGKI and hypertension seems to depend on the pathogenesis of the disease.

Nevertheless, a reduced expression of cGKI has been described in neointimal tissue of patients undergoing coronary angioplasty revascularization [32], suggesting that the enzyme can be downregulated in different pathological conditions. Therefore, further research on cGKI distribution and activity in humans will contribute to a better understanding of cardiovascular diseases and focus on the cGKI gene as a candidate for new gene target therapy.

**Acknowledgements**

This work is dedicated to the memory of Prof. Alberto Giotti. The study was supported by grants from the University of Florence.

**Disclosure Statement**

No conflicts of interest are declared by the authors.

---

**References**


