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The heart initially compensates for hypertension-mediated pressure overload by enhancing its contractile force and developing hypertrophy without dilation. G_q protein–coupled receptor pathways become activated and can depress function, leading to cardiac failure. Initial adaptation mechanisms to reduce cardiac damage during such stimulation remain largely unknown. Here we have shown that this initial adaptation requires regulator of G protein signaling 2 (RGS2). Mice lacking RGS2 had a normal basal cardiac phenotype, yet responded rapidly to pressure overload, with increased myocardial G_q signaling, marked cardiac hypertrophy and failure, and early mortality. Swimming exercise, which is not accompanied by G_q activation, induced a normal cardiac response, while Rgs2 deletion in G_\alpha_q-overexpressing hearts exacerbated hypertrophy and dilation. In vascular smooth muscle, RGS2 is activated by cGMP-dependent protein kinase (PKG), suppressing G_q-stimulated vascular contraction. In normal mice, but not Rgs2^{−/−} mice, PKG activation by the chronic inhibition of cGMP-selective phosphodiesterase 5 (PDE5) suppressed maladaptive cardiac hypertrophy, inhibiting G_q-coupled stimuli. Importantly, PKG was similarly activated by PDE5 inhibition in myocardium from both genotypes, but PKG plasma membrane translocation was more transient in Rgs2^{−/−} myocytes than in controls and was unaffected by PDE5 inhibition. Thus, RGS2 is required for early myocardial compensation to pressure overload and mediates the initial antihypertrophic and cardioprotective effects of PDE5 inhibitors.

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Regulator of G protein signaling 2 mediates cardiac compensation to pressure overload and antihypertrophic effects of PDE5 inhibition in mice

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The heart initially compensates for hypertension-mediated pressure overload by enhancing its contractile force and developing hypertrophy without dilatation. G α-protein–coupled receptor pathways become activated and can depress function, leading to cardiac failure. Initial adaptation mechanisms to reduce cardiac damage during such stimulation remain largely unknown. Here we have shown that this initial adaptation requires regulator of G protein signaling 2 (RGS2). Mice lacking RGS2 had a normal basal cardiac phenotype, yet responded rapidly to pressure overload, with increased myocardial G q signaling, marked cardiac hypertrophy and failure, and early mortality. Swimming exercise, which is not accompanied by G α activation, induced a normal cardiac response, while Rgs2 deletion in G q overexpressing hearts exacerbated hypertrophy and dilatation. In vascular smooth muscle, RGS2 is activated by cGMP-dependent protein kinase (PKG), suppressing G α-stimulated vascular contraction. In normal mice, but not Rgs2–/– mice, PKG activation by the chronic inhibition of cGMP-selective phosphodiesterase 5 (PDE5) suppressed maladaptive cardiac hypertrophy, inhibiting G α q-coupled stimuli. Importantly, PKG was similarly activated by PDE5 inhibition in myocardium from both genotypes, but PKG plasma membrane translocation was more transient in Rgs2–/– myocytes than in controls and was unaffected by PDE5 inhibition. Thus, RGS2 is required for early myocardial compensation to pressure overload and mediates the initial antihypertrophic and cardioprotective effects of PDE5 inhibitors.

Introduction
The adult heart responds to sustained pressure overload by developing ventricular hypertrophy. The signaling events mediating this process are substantially driven by activation of GPCRs, which in turn stimulate multiple downstream intracellular cascades. Exposure to sufficient magnitude and duration of GPCR stimulation is thought to be necessary to tip the balance between adaptive and maladaptive responses (1, 2). However, the response may also depend upon how well the heart can mount countermeasures to effectively blunt such adverse signaling.

One set of negative controllers of GPCRs is the family of more than 30 regulator of G protein signaling (RGS) proteins (3). Upon GPCR activation, GDP is exchanged for GTP on the G α subunit, allowing for dissociation from G βγ subunits and activation of downstream effectors. RGS proteins inhibit these cascades by accelerating G α-dependent GTP hydrolysis to reconstitute the heterotrimeric G protein complex. RGS proteins also act as effector antagonists by physically blocking the binding of G protein subunits to their protein targets and interfering with downstream signaling proteins (4). RGS2–RGS5 are thought to be important in the heart (5, 6), although their precise roles remain unclear. Human heart failure is associated with increased RGS4 expression, whereas expression of RGS2 is unchanged (7). Forced overexpression of RGS4 blunts G α q-stimulated cardiac hypertrophy in rat neonatal cardiac myocytes (8) and intact hearts (9) and suppresses cardiac hypertrophy of transgenic mice lacking guanylate cyclase-A (natriuretic peptide-stimulated cyclase; ref. 10). Yet other studies in hearts overexpressing RGS4 found rapid cardiac dilatation and marked mortality upon pressure overload induction (11), highlighting a complex role. This was further demonstrated by the recent discovery that RGS4 regulates parasympathetic (G α<sub>qi/oq</sub>) signaling to control heart rate in the sinoatrial node (12).

In contrast to RGS4, which inactivates multiple G α proteins (13), myocyte RGS2 appears more selective for G α q (6, 14). Given the recognized and prominent role of G α q signaling to maladaptive remodeling as a result of pressure overload (15, 16), RGS2 is an intriguing candidate as an intrinsic suppressor of this pathobiology. Mice globally lacking RGS2 were found to develop modest systemic hypertension, although they exhibit no major cardiac phenotype (17), which suggested that RGS2 had a modest role in the heart. Yet more recent studies found that knockdown of the gene encoding RGS2 amplifies hypertrophic responses in neonatal myocytes exposed to G α q stimuli (18).

Nonstandard abbreviations used: CaMKII, Ca<sup>2+</sup>-calmodulin–dependent kinase II; Cn, calcineurin; ET1, endothelin-1; FS, fractional shortening; PDE5, phosphodiesterase 5; PLC, phospholipase C; PV, pressure-volume; RGS, regulator of G protein signaling; TAC, transverse aortic constriction.

Conflict of interest: The authors have declared that no conflict of interest exists.

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An important feature of both RGS2 and RGS4 is that they are activated by PKG, attenuating Gq-coupled vasoconstriction in vascular (19) and gastric (20) smooth muscle, and, in the case of RGS4, enhancing antihypertrophic effects of natriuretic peptides (10). Even more pronounced suppression of cardiac hypertrophy coupled to PKG activation has been achieved by inhibiting phosphodiesterase 5 (PDE5; ref. 21) with drugs widely used to treat erectile dysfunction (e.g., sildenafil). Given its greater selectivity for Gq, we hypothesized that RGS2 plays a particularly central role in the antihypertrophic effects of this therapy. Understanding such mechanisms has taken on clinical relevance, given the recently initiated NIH multicenter trial of the PDE5 inhibitor sildenafil for treating heart failure with a normal ejection fraction (RELAX study; http://clinicaltrials.gov/ct2/show/NCT00763867).

In the present study, we tested the role of RGS2 in pressure overload remodeling and its amelioration by PDE5 inhibition. Mice genetically lacking RGS2 did not compensate against pressure overload, but instead developed rapid and marked cardiac hypertrophy and dysfunction and early lethality. In contrast, their response to swimming exercise was normal. In addition to targeting myocardial Gq signaling, RGS2 was found to be essential in order for PDE5 inhibition to reduce hypertrophy and improve cardiac function upon exposure to pressure overload.

**Results**

RGS2 is expressed in adult mouse myocytes and suppresses Gq stimulation. RGS2 protein expression was detected in isolated adult myocytes from control Rgs2+/+, but not Rgs2−/−, mouse hearts (Figure 1A). RGS2-mediated protection against myocyte hypertrophy has been previously reported in neonatal myocytes using RGS2 RNAi (18). Here, we examined adult myocytes from Rgs2+/+ and Rgs2−/− mouse hearts and found enhanced protein synthesis in cells incubated with the Gq agonist endothelin-1 (ET1). This finding supports the role of RGS2 in countering such stimulation in adult cells.

Rgs2−/− mouse hearts develop marked hypertrophy and failure and early lethality from pressure overload. To test whether RGS2 is cardioprotective against development of hypertrophy in vivo, Rgs2−/− mice (4–5 months old) were subjected to pressure overload by transverse aortic constriction (TAC; ref. 21). Resting cardiac anatomy and global function, as assessed by echocardiography, was similar between Rgs2−/− and littermate control Rgs2+/+ mice (Supplemental Table 1; supplemental material available online with this article; doi:10.1172/JCI35620DS1). After TAC, however, Rgs2−/− mice developed marked hypertrophy (Figure 1C) and had a high mortality rate within 1 wk, whereas all Rgs2+/+ mice survived this period (Figure 1D). Marked lung congestion (wet lung weight) was documented in surviving and deceased Rgs2−/− animals (Figure 1E), which supported heart failure as the cause of death. The increased heart mass was caused by both exacerbated myocyte hypertrophy and fibrosis (Figure 1F). Cardiac apoptosis was not observed within this 1 wk time period in either genotype (data not shown). TAC-induced changes in cardiac fetal genes Nppa, Nppb, and Myb7 were greater in Rgs2−/− than in Rgs2+/+ hearts, whereas expression of Atp2a2 declined in both models similarly (Supplemental Figure 1).

Chamber dilation and lack of functional compensation in Rgs2−/− TAC mouse hearts. Rgs2−/− hearts dilated after pressure overload, with end-diastolic and end-systolic dimensions increasing and fractional shortening (FS) declining more than in Rgs2+/+ hearts (Figure 2A). These echocardiographic data were supported by more detailed invasive pressure-volume (PV) analysis (Figure 2, B–D). At rest, Rgs2−/− mice had LV peak systolic pressure that was increased about 15 mmHg compared with Rgs2+/+ mice and increased total LV afterload, as measured by effective arterial elastance (Figure 2C). However, basal cardiac systolic and diastolic function was essentially identical between the groups (Supplemental Table 2). TAC increased cardiac afterload to near-identical levels in both genotypes (Figure 2C), because load largely depended on the proximal aortic constriction. Similarities of loading changes after 48 h TAC were also confirmed (Supplemental Figure 2). In Rgs2−/− mice, 1 wk TAC triggered functional compensation with enhanced contractility: the end-systolic PV relation shifted leftward with increased slope (Figure 2B). This did not occur in Rgs2+/+ mice; instead, hearts dilated with a rightward shift of PV relations (Figure 2B). Summary data for contractility (end-systolic PV relation slope, volume position, peak rate of LV pressure rise, and preload recruitable stroke work) and relaxation (peak rate of LV pressure decline and time constant tau) are provided in Figure 2D. Unlike control hearts, Rgs2−/− hearts failed to compensate by increasing contractility and had impaired relaxation, with net cardiac output depending more on marked chamber dilation (Frank-Starling law).

Early and robust activation of Gq-related pathways in Rgs2−/− TAC mouse hearts. Because RGS2 principally suppresses Gq signaling, we examined the activation of the phosphatase calcineurin (Cn), Ca2+-calmodulin–dependent kinase II (CaMKII), and the MAPKs ERK1/2, JNK, and p38, all of which are potentially activated by Gq-coupled stimuli and are associated with hypertrophy and/or cardiac dilation (2, 22). After 1 wk TAC, Cn protein expression rose modestly in Rgs2+/+ hearts, but markedly increased in Rgs2−/− hearts (Figure 3A). Parallel changes were observed in the gene expression of regulator of Cn-1 (RCAN-1), an index of Cn activity (Figure 3B). This was observed even after 48 h TAC in Rgs2−/− hearts, a time at which expression was otherwise unaltered. CaMKII activation also markedly increased after 48 h TAC in Rgs2−/− hearts and persisted at 1 wk TAC, whereas TAC Rgs2−/− hearts showed minimal CaMKII activation (Figure 3C). Among the 3 MAPKs, we observed greater activation of ERK1/2 and JNK at both time points in Rgs2−/− hearts (Figure 3D). Phosphorylation of p38 rose markedly and similarly in both groups after 48 h TAC, but declined at 1 wk TAC, which suggests that it was regulated by alternative cascades. These early signaling responses were not coupled to changes in RGS2 expression in control myocardium or myocytes (data not shown).

Gq is a critical target of RGS2. To test whether Gq is indeed the critical target of RGS2 in the heart, we performed several studies whereby Gq signaling was modified. Activation of phospholipase Cβ (PLCβ) is a primary mechanism for Gq signaling and is required for dilated cardiomyopathy to develop in cardiac-targeted Gαs-overexpressing transgenic animals (referred to herein as Gαs-OE; ref. 23). We therefore subjected Rgs2−/− hearts to 48 h TAC with or without cotreatment by a PLCβ inhibitor (U73122, 2.5mg/kg/d) or inactive inhibitor, whereas Gq signaling was modified. Activation of phospholipase Cβ (PLCβ) is a primary mechanism for Gq signaling and is required for dilated cardiomyopathy to develop in cardiac-targeted Gαs-overexpressing transgenic animals (referred to herein as Gαs-OE; ref. 23). We therefore subjected Rgs2−/− hearts to 48 h TAC with or without cotreatment by a PLCβ inhibitor (U73122, 2.5mg/kg/d) or inactive inhibitor, whereas the inactive analog at the same dose had no impact (Figure 4, A and B; see Supplemental Figure 2B for summary echo data).

In a second series of experiments, Gαs-OE mice (24) were bred with Rgs2−/− animals, and cardiac function and morphology were examined at 3–4 wk of age. Gαs-OE/Rgs2−/− mice exhibited minimal hypertrophy, but slight chamber dilation and reduced FS. However, these 3 were substantially worsened in Gαs-OE/Rgs2+/+ mice (Figure 4, C–E).
Figure 1
Cardiac phenotype of Rgs2−/− mice. (A) Myocyte RGS2 protein expression was detected in Rgs2+/+ (+/+ cells) but not Rgs2−/− (−/−) cells. (B) Adult mouse myocytes (n = 5 per group) from Rgs2−/− hearts displayed an amplified growth response to ET1, as assessed by radiolabeled leucine incorporation. *P value shown is for interaction of genotype and condition (2-way ANOVA). *P < 0.01; **P < 0.001. (C) Representative whole hearts, H&E-stained cross sections, and summary results for heart weight normalized to tibia length (HW/TL) in Rgs2+/+ and Rgs2−/− hearts subjected to 1 wk TAC. White bar shows data from deceased mice. (D) Kaplan-Meier survival curve showing markedly increased mortality in Rgs2−/− mice subjected to TAC compared with littermate controls. (E) Wet lung weight normalized to body weight (n = 8–13 per group). White bar shows data from deceased mice. (F) Representative Masson’s trichrome staining of the heart section. Blue stain indicates collagen deposition. Scale bars: 100 μm. Also shown are summary quantification results on myocyte diameter and collagen fraction (n = 4 hearts per group; >50 cells per heart; 5–6 sections for fibrosis analysis). *P < 0.05 versus Rgs2+/+ sham; §P < 0.05 versus Rgs2−/− sham; †P < 0.05 versus Rgs2+/+ TAC.
Overexpression of $G_{\alpha q}$ itself lowered heart rate, as described previously (24), but this was not altered by deleting Rgs2.

Finally, we examined the cardiac response to a stress that did not involve $G_{\alpha q}$ stimulation. $Rgs2^{-/-}$ and $Rgs2^{+/+}$ mice were subjected to 6 wk of swimming exercise (twice daily for 90 min). This regimen stimulated an increase in LV mass of about 30% in both groups, with no impairment of cardiac function (Figure 4F). Protein analysis confirmed that Cn, CaMKII, and ERK1/2 were not activated in either genotype by the exertional stress (Supplemental Figure 3). Thus, unlike our findings with TAC or

Figure 2

$Rgs2^{-/-}$ hearts display chamber dilation and fail to compensate to TAC. (A) Echocardiographic data before and after 1 wk TAC ($n = 7$ per group). LV-Dd, LV end-diastolic dimension; LV-Ds, LV end-systolic dimension. $^aP < 0.05$ versus $Rgs2^{+/+}$ 1 wk TAC. (B) Representative PV loops during preload reduction by inferior vena cava occlusion in sham (black line) and 1 wk TAC animals (thin dotted line). Steepness of left upper relation (end-systolic elastance [Ees]) reflected contractile function and was enhanced after TAC in $Rgs2^{-/-}$ mice, but unaltered with a right-shift (remodeling) of the relation in $Rgs2^{+/+}$ mice. (C) Peak systolic LV pressure (LVP sys) and effective arterial elastance (Ea; an index of total ventricular afterload). $Rgs2^{-/-}$ mice had somewhat higher basal afterload, but both genotypes had similarly increased afterload after 1 wk TAC ($n = 5$–7 per group). $P$ value shown is for interaction of genotype and condition. $^aP < 0.05$ versus sham; $^bP < 0.05$ versus $Rgs2^{+/+}$ sham. (D) Summary data obtained from PV loop analysis shown in bar graphs. $P$ values shown are for interaction of genotype and condition (2-way ANOVA). $^aP < 0.001$, $^bP < 0.05$ versus sham. $V_{100}$, volume position (end-systolic volume at common end-systolic pressure — 100 mmHg — derived from end-systolic PV relation); $dP/dt_{\text{max}}$, peak rate of LV pressure rise; PRSW, preload recruitable stroke work; $dP/dt_{\text{min}}$, peak rate of LV pressure decline; Tau, relaxation time constant; SV, stroke volume; CO, cardiac output. $P$ values shown are for interaction of genotype and condition (2-way ANOVA). $^aP < 0.05$, $^bP = 0.06$ versus sham; $^cP < 0.05$ versus corresponding $Rgs2^{-/-}$ TAC and $Rgs2^{-/-}$ sham (1-way ANOVA).
G_{aq}-OE^- animals, hypertrophic response that did not involve G_q signaling stimulation was not exacerbated by a lack of RGS2.

Antihypertrophic effect of PDE5 inhibition is absent in Rgs2^{−/−} mouse hearts. In noncardiac cells, RGS2 is activated by PKG-1 binding and phosphorylation, inducing their translocation to the outer plasma membrane to inactivate G_q (19). In hearts exposed to stress, cardiac PKG can be potently activated by inhibiting the cGMP hydrolytic enzyme PDE5, which in turn suppresses hypertrophy and/or dys-function in mice subjected to pressure overload (21). While various individual PKG targets may underlie this response (25), RGS2 could reflect a very proximal effector that subsequently regulates multiple downstream cascades. To test the importance of RGS2 to this regulation, mice were fed with the PDE5 inhibitor sildenafil during 1 wk TAC. Sildenafil blunted hypertrophy in controls, reducing wall thickness and chamber size and improving FS, but had no effect on these properties in Rgs2^{−/−} TAC hearts (Figure 5, A–C).
Sildenafil also improved contractility and relaxation in Rgs2+/+ mice exposed to TAC, which was not observed in Rgs2–/– mice (Fig. 5, D and F). As previously reported (21), sildenafil did not itself reduce cardiac afterload (Figure 5E).

Because Rgs2–/– mice displayed greater activation of Gq-linked pathways (Cn, CaMKII, and ERK1/2), we tested whether sildenafil suppresses these signaling pathways in Rgs2+/+ and Rgs2–/– hearts. Activation of each enzyme was indeed blunted by sildenafil cotreatment in Rgs2+/+ TAC hearts, an effect not observed in Rgs2–/– TAC hearts (Figure 5E).

Figure 4
Gqα as a critical target of RGS2. (A) Inhibition of PLCβ with U73122 prevents exacerbated hypertrophy (heart weight/tibia length), chamber dilation, and dysfunction (n = 7–14 per group). *P < 0.001 versus other groups. (B) In Rgs2–/– mice subjected to 48 h TAC, the inactive control agent U73343 did not suppress pathological remodeling. Summary echo data are provided in Supplemental Figure 2B. (C) Cardiac hypertrophy was exacerbated in double-mutant GqαOE–Rgs2–/– mice (n = 7–9 per group). **P < 0.01 versus other groups. (D) Corresponding echocardiograms showed worsened function and chamber dilation. (E) Summary data for LV diastolic dimension, FS, and heart rate (n = 7–9 per group). *P < 0.05 versus other groups; †P < 0.05 versus all GqαOE– groups. (F) Response of cardiac LV mass and echocardiographic FS in mice subjected to swimming versus sedentary animals. Exercise-induced hypertrophy was similar in Rgs2+/+ and Rgs2–/– mice, with no change in FS. *P < 0.01, †P < 0.05 versus respective week-0 baseline.
substantial further enhancement with sildenafil treatment (Figure 7A; $P = 0.37$, genotype and genotype × condition; $P < 0.0001$, condition effect; 2-way ANOVA). Thus, the lack of a sildenafil effect in $\text{Rgs2}^{-/-}$ could not be attributed to lack of PKG activation per se.

PKG-1α activates RGS2 in vascular smooth muscle (19), leading to translocation of both proteins to the outer cell membrane; furthermore, PKG-1α is the prominent isoform in the heart (25). Therefore, we examined the localization of both proteins in cardiac myocytes. In control cells, RGS2 displayed faint diffuse cytosolic and membrane localization, whereas PKG-1α was diffusely distributed with a somewhat striated pattern (Figure 7B). After 2 h exposure to the $G_q$ agonist ET1, both RGS2 and PKG-1α intensified at the outer membrane (Figure 7B). Identical results were obtained using angiotensin II stimulation (data not shown). This translocation was fully blocked by coinubcation with the specific PKG peptide inhibitor DT2 (26) and was conversely stimulated by 8Br-cGMP.

Figure 5
Amelioration of pressure load–induced cardiac hypertrophy/remodeling via PDE5 inhibition with sildenafil is absent in $\text{Rgs2}^{-/-}$ mice. (A) Sildenafil (Sil) suppressed hypertrophy in $\text{Rgs2}^{+/+}$, but not $\text{Rgs2}^{-/-}$, mice ($n = 8–13$ per group). (B and C) Representative echocardiograms and summary data showing response to 1 wk TAC (T) with or without sildenafil treatment ($n = 7–9$ per group). S, sham. (D) Effects of sildenafil on LV PV loops. (E) Sildenafil did not alter the increase in LV afterload in either genotype ($n = 4–7$ per group). (F) Summary invasive PV loop results ($n = 4–7$ per group). Sildenafil improved systolic and diastolic function in $\text{Rgs2}^{+/+}$ TAC hearts, but had no effect in $\text{Rgs2}^{-/-}$ TAC hearts. *$P < 0.05$ versus respective sham control; †$P < 0.05$ versus respective 1 wk TAC; ‡$P < 0.05$ versus $\text{Rgs2}^{+/+}$ TAC; §$P < 0.05$ versus $\text{Rgs2}^{-/-}$ sham.
Importantly, translocation of both proteins was also observed after TAC in Rgs2+/+ hearts, evident at 48 h and declining by 1 wk. Sildenafil treatment enhanced membrane localization of both proteins after 1 wk TAC, consistent with enhanced PKG activity. In Rgs2−/− cells, basal PKG-1α localization was similar to Rgs2+/+ cells (Supplemental Figure 4A). Intriguingly, PKG-1α also translocated to the outer membrane after 48 h TAC; however, this was mostly absent by 1 wk TAC (Figure 7C). Moreover, unlike controls, concomitant inhibition of PDE5 did not significantly alter this localization. These confocal results were confirmed by cell-fraction protein immunoblots (Figure 7D; gel loading data provided in Supplemental Figure 4B).

**Discussion**

The present study demonstrates a major cardiac regulatory role for RGS2 as a critical brake against the Gαq/PLCβ pathway in the early cardiac stress response to pressure overload. Without RGS2, pressure overload induced rapidly exacerbated hypertrophy and dysfunction, amplified Gαq-coupled signaling, and early mortality.

Deletion of Rgs2 in Gαq transgenic hearts worsened dilation and hypertrophy, whereas inhibiting PLCβ blocked the rapid deterioration and hypertrophy after TAC in Rgs2−/− hearts. Swimming, a non-Gαq-coupled stress (1, 27), had no adverse effect in these mice. Finally, we showed that the capacity of cGMP/PKG stimulated by PDE5 inhibition to blunt hypertrophy and enhance cardiac function within the first week of pressure overload stress was critically coupled to RGS2 and its suppression of Gαq signaling. Rgs2−/− mice exhibited a very rapid hypertrophic response to pressure overload, revealing both the early potency of Gαq stimulation and a key role of its suppression by RGS2 to permit early compensation. This finding demonstrates that the transition from adaptive to maladaptive hypertrophy involves not only the magnitude and duration of proremodeling stimuli, but also the robustness and sustainability of critical countermeasures that blunt their toxicity. Stimulation of Gαq by TAC can reflect both receptor/agonist interaction from neurohormones and mechanical stretch, as described previously via the angiotensin II receptor (28, 29).

The Rgs2−/− phenotype is somewhat unusual in that animals have

**Figure 6**

Sildenafil suppresses Gαq-coupled molecular cascades in Rgs2+/+ but not Rgs2−/− hearts. (A) Protein expression for Cn. (B and C) Phosphorylated and total protein expression for CaMKII (B) and ERK1/2 (C). Summary data are shown for each (n = 3–6 per group). †P < 0.05 versus Rgs2+/+ 1 wk TAC.
Figure 7
Regulation of RGS2 by PKG-1α. (A) PKG is similarly activated with and without concomitant sildenafil treatment in both genotypes. *P < 0.05 versus respective sham; †P < 0.005 versus TAC alone. (B) Subcellular localization of RGS2 and PKG-1α in Rgs2+/+ mouse adult cardiac myocytes by immunohistochemistry using confocal microscope. Images represent baseline (control), ET1 stimulation, ET1 stimulation with PKG inhibitor (ET1+DT2), and PKG stimulation with 8Br-cGMP. More than 20 cells were analyzed from 1 experiment, and experiments were repeated 3 times. Original magnification, ×200. (C) Translocation of PKG-1α to outer membrane with 48 h TAC occurred in both genotypes and declined somewhat after 1 wk in Rgs2+/+ mice but was virtually absent in Rgs2−/− mice at the same time. Sildenafil treatment restored membrane localization and enhanced RGS2 membrane signal in Rgs2+/+ cells, but had no impact in Rgs2−/− cells. Original magnification, ×200. (D) Immunoblot of PKG-1α from cytosolic (soluble [S]) and membrane (particulate [P]) fractions in sham control, 48 h, and 1 wk TAC myocardium. Summary data of particulate/soluble ratio is shown at right (n = 3–6 per group). *P < 0.001 versus sham; †P < 0.001 versus 1 wk TAC; ‡P < 0.001 versus 48 h TAC. Equal protein loading was confirmed by Ponceau staining (Supplemental Figure 4B).
normal heart function and LV mass at rest, yet develop marked hypertrophy, dilation, and early mortality after TAC. Genetic models lacking cardiac protective or structural genes that exhibit early mortality after TAC (e.g., gp130 and integrin β1-interacting protein; refs. 30, 31) dilate without hypertrophy, whereas others (e.g., dominant-negative thiorodoxin-1 and inactivated glycogen synthase kinase-3β; refs. 32, 33) induce hypertrophy that worsens further with TAC, yet do not display early lethality.

Other RGS proteins, such as RGS3 and RGS4, may also be involved in hypertrophy and failure regulation (10, 34), although these proteins are less specific to Gq (6). Mice with myocyte-targeted RGS4 overexpression exhibit a complex phenotype, developing rapid dilation and early mortality (1–2 days) without hypertrophy after exposure to TAC, but survivors exhibit less eventual hypertrophy (11). When these animals are genetically crossed with GαqOE mice, cardiac abnormalities in the latter strain are ameliorated (9).

Thus, RGS4 may have protective roles in the later stages of disease, consistent with its upregulation in end-stage human heart failure (7, 35). RGS4 may also mediate ameliorative effects of natriuretic peptides on cardiac stress remodeling, although in vivo support for this hypothesis is so far based on the ability of RGS4 overexpression to blunt the cardiac phenotype in mice genetically lacking natriuretic peptide receptor–coupled guanylate cyclase–A (10).

RGS2 modulates stress response signaling by several mechanisms (36, 37). First, it inactivates the Gαq subunit, which can suppress downstream Ca2+-activated cascades (including Cn and CaMKII) and MAPKs (including ERK and JNK), as shown in the current study. Cn and CaMKII in particular are considered potent contributors to maladaptive remodeling (22, 38). The results of the present studies using PLCβ inhibition, genetic deletion of Rgs2 in GαqOE mice, and swimming exercise support RGS2 suppression of Gαq/PLCβ signaling as a central feature of its cardioprotective effects. RGS proteins can also inhibit Gβγ-induced signaling by reforming the G protein heterotrimer, and this could influence Akt/glycogen synthase kinase–3β cascades (39), which we have previously shown to be suppressed by PDE5 inhibition (21). RGS proteins can further bind to effector enzymes, such as adenylate cyclase and PKC isoforms, to blunt signaling (40), although the role of this modulation in heart remains unknown.

While RGS2 has high affinity for Gq, it can suppress Gqβγ or Gq in some cell types. For example, it regulates presynaptic Ca2+ channels via Gqβγ subunits in neurons (41) and impacts carbachol-stimulated activation of ERK and Akt in COS cells via Gqβγ interaction (42). RGS2 blunts Gq/coupled cAMP accumulation triggered by parathyroid hormone–related peptide (43). However, the effects of RGS2 over Gqβγ or Gq appear to be unimportant in adult cardiac myocytes, as neither muscarinic nor β-adrenergic CAMP regulation is altered by RGS2 overexpression in contrast to RGSS3, RGSS4, or RGSS5 (6).

Even though PDE5 is more abundantly expressed in vascular smooth muscle, antihypertrophic effects from its inhibition are likely caused by direct cardiac effects. First, as demonstrated in the present study, cardiac afterload associated with TAC was unchanged by sildenafil treatment. Second, PDE5 inhibition suppresses Cn/nuclear factor of activated T cell (Cn/NFAT) signaling and expression of the genes encoding atrial and brain natriuretic peptides in isolated myocytes stimulated with a Gq agonist (21, 44). Third, knockdown of the gene encoding PDE5 in cardiac myocytes is antihypertrophic, having an impact similar to that of sildenafil, and the effect of combining both is similar to that of each alone (44). This last finding supports specificity of sildenafil for inhibiting PDE5. The improvement in cardiac function by sildenafil involves several mechanisms, including enhanced myocyte calcium cycling and a decline in PKCα activation (45), both potentially linked to RGSS2 suppression of Gq/PLCβ.

Growing evidence supports an important role for myocardial cGMP/PKG signaling as an intrinsic brake to suppress maladaptive cardiac stress remodeling (46); however, only recently have studies revealed potential mechanisms for this effect. The first reported target was Cn/NFAT, which can be suppressed in neonatal myocytes by enhancing cGMP (e.g., NO donors) or expressing activated PKG (47). PDE5 inhibition was shown to have a similar effect in vivo (21), and our present data further support this. PKG also blunts RhoA activation by phosphorylation at Ser188 (48), although whether this applies to myocytes remains unknown. PKG phosphorylates SMAD3 at Ser309 and Thr388 in fibroblasts, preventing its nuclear translocation to block TGF-β1 activation (49). This may be important for how PDE5 inhibitors suppress fibrosis. PKG can also block p38 autophosphorylation coupled to its binding to the scaffold protein TGF-β–activated protein kinase 1–binding protein 1 (TAB1) in ischemia/reperfusion injury (50), although this mechanism does not appear to be involved in 1 wk TAC. While the latter pathways involve direct PKG phosphorylation, similar mechanisms have not been identified for Cn/NFAT or CaMKII suppression. However, as these enzymes are coupled to Gq stimulation, suppression of the proximal signal by RGS2-PKG activation could provide a mechanism as well as explain reduced RhoA and TGF-β stimulation.

In vascular smooth muscle cells, activation of RGS2 by the nitric oxide/cGMP pathway requires binding to PKG-1α and its phosphorylation at Ser46 and Ser64 by PKG-1α, but RGS2 does not need the Gq-coupled pathway to be activated (19). cGMP stimulation also increases the association of the smooth muscle cell plasma membrane with wild-type but not mutant (S46,64A) RGS2 (51), highlighting the importance of PKG activity for this association and GPCR modulation. The current study is the first to our knowledge to show that PKG-1α rapidly translocates to the outer plasma membrane from a resting cytosolic distribution in adult mouse myocytes in vitro and in vivo, concurrent with RGS2 plasma membrane localization, and that PKG activation is required for this movement. The acute movement of PKG-1α to the plasma membrane even in Rgs2–/– cells indicates that another chaperone is involved, if not more than one. This may require binding to the leucine zipper motif — translocates the enzyme to the nucleus upon activation in BHK cells (52). Importantly, PKG-1α translocation was more transient in Rgs2–/– myocytes than in controls and did not provide anti-hypertrophic protection. Rgs2–/– TAC hearts showed relative PKG activation by 1 wk sildenafil treatment similar to that of Rgs2+/- hearts; thus, compartmentalization of this signal and its specific interaction with RGS2 appears to be central to its capacity to suppress Gq signaling and maladaptive hypertrophy.

Our study has several limitations. The Rgs2–/– mice globally lacked the gene, and though cardiac function was unchanged (observed even in 10-month-old animals; Supplemental Table 3), adaptive responses to sustained modest hypertension could still occur. Although there were no basal differences in the signaling cascades examined, our analysis was admittedly quite focused. Importantly, TAC-induced pressure overload was similar in both genotypes, and whatever adaptations may have existed, they failed...
to compensate to the higher afterload. The blood pressure disparity we observed was somewhat less than originally reported (17), yet similar to other studies using the same mouse model (53). Some of this disparity may relate to the anesthesia used in our protocol. Because the gene was absent from conception, we cannot yet assess the role of RGS2 in the later stages of pressure overload disease. Such assessment will require an inducible gene deletion model currently under development. However, our present results indicate that a dominating PKG effector in the initial response to pressure overload is RGS2.

In summary, our study demonstrates an essential role for RGS2—inhibiting the Gαq/PLCβ pathway—in early compensatory hypertrophy development to pressure overload and provides a mechanism for antihypertrophic effects associated with PKG enhancement from PDE5 inhibition. This coactivation provides what we believe to be the most proximal target for PKG-mediated inactivation of hypertrophy stimulation thus far described and can explain how multiple distal pathways can be concomitantly suppressed by PDE5 inhibition.

Methods

Animal models. Rgs2−/− mice, which harbor a global RGS2 deletion, and Rgs2+/− littermate controls in C57BL/6 background were as reported previously (19), originally developed by J. Penninger (54). Gαq−/−OE mice (40 copies of the transgene, FVB background; ref. 24), generously provided by G. Dorn (Washington University Center for Pharmacogenomics, St. Louis, Missouri, USA), were crossed with Rgs2−/− animals to generate animals lacking Rgs2 and overexpressing Gαq in the heart. The same generation of animals (Gαq−/−OE Rgs2−/−, Gαq−/−OE Rgs2+/−, Gαq−/−OE Rgs2−/−, Gαq−/−OE Rgs2−/−) was also studied.

TAC. Pressure overload was produced by TAC in male Rgs2−/− and Rgs2+/− animals of 4–5 months of age (mean BW, 26 g) as previously described (21). Briefly, animals were anesthetized with isoflurane (2%–3%), intubated, and mechanically ventilated. The transverse aorta was constricted with a 26-gauge needle using 7-0 prolene suture, after which the chest was closed and the animal was allowed to recover from anesthesia. Control mice were subjected to sham operations, and animals were studied 48 h to 1 wk after surgery. Sets of male Rgs2−/− animals were subjected to 48 h TAC and treated daily with 2.5 mg/kg/d U73122, a specific PLCβ inhibitor, or U73343, its inactive analog, in peanut oil via i.p. injection. Additional animals in either genotype were subjected to 1 wk TAC while cotreated with sildenafil (200 mg/kg/d) mixed in food (Bioserve soft diet).

Swimming exercise. Male Rgs2−/− and Rgs2+/− animals (3 months of age; 23–25 g BW) were studied. The forced swimming program was a modified 6-wk protocol based on a prior method (55). Control animals remained sedentary in a cage for 6 wk. Briefly, 10 mice at a time swam twice a day in a 45-cm by 75-cm container with the water kept at approximately 32°C. Sessions began at 10 min and were increased by 10-min increments each day until each session lasted 90 min; this was then maintained for the remainder of the 6-wk period. Constant monitoring ensured the safety of the mice and prevented them from floating or holding their breath under water. All animal protocols were approved by the Animal Care and Use Committee of Johns Hopkins University.

Echocardiography. In vivo cardiac morphology was assessed by transthoracic echocardiography (Acuson Sequoia C256, 13-MHz transducer; Siemens) in conscious mice (21). M-mode LV end-systolic and end-diastolic dimensions were averaged from 3–5 beats. LV percent FS and LV mass were calculated as described previously (21). Wall thickness of lateral free wall and intraventricular septum were averaged. Studies and analysis were performed by investigators blinded to genotype or heart condition.

In vivo hemodynamics. In vivo LV function was assessed by PV catheter as described previously (21). Briefly, mice were anesthetized with 1%–2% isoflurane, 750–100 mg/kg urethane i.p., 5–10 mg/kg etomidate i.p., and 1–2 mg/kg morphine i.p.; were subjected to tracheostomy; and were ventilated with 6–7 μl/g tidal volume and 130 breaths/min. Volume expansion (12.5% human albumin, 50–100 μl over 5 min) was provided through a 30-gauge cannula via the right external jugular vein. The LV apex was exposed through an incision between the seventh and eighth ribs, and a 1.4-Fr PV catheter (SPR 839; Millar Instruments Inc.) was advanced through the apex to lie along the longitudinal axis. Absolute volume was calibrated, and PV data were measured at steady state and during transient reduction of venous return by occluding the inferior vena cava with a 6-0 silk snare suture. Data were digitized at 2 kHz, stored to disk, and analyzed with custom software. From the 10–15 successive cardiac cycles during the inferior vena cava occlusion, the end-systolic PV relation slope (i.e., end-systolic elastance) and stroke work–end-diastolic volume relation (i.e., preload recruitable stroke work) were derived.

Tissue histology. Myocardium fixed with 10% formalin was analyzed for myocyte hypertrophy and fibrosis. Tissue was paraffin embedded, cross-sectioned into 5–8-μm slices, and stained with H&E or Masson’s trichrome staining (21). We analyzed 6 serial sections of mid-LV per heart. To assess mean cardiomyocyte diameter and interstitial collagen fraction, 6–8 regions of photomicrographs covering the whole section were obtained and quantified using computer-assisted image analysis (Adobe Photoshop version 5.0 and NIH Image J). Average data reflect results from 4–5 hearts per group (>50 cells).

Western blot analysis. Protein was prepared from snap-frozen heart tissue or isolated cardiac myocytes, and fractionation was performed as previously described (21, 56). After homogenization in 25 mM Tris-HCL, pH 7.5, 4 mM EGTA, 2 mM EDTA, 5 mM dithiothreitol, 1 mM phenylmethylsulfonyl fluoride, and 1 μM leupeptin and incubation on ice for 30 min, samples were spun at 100,000 g for 30 min at 4°C. The supernatant was saved as the cytosolic fraction, and the pellet was resuspended in homogenization buffer with 1% Triton X-100 added. This was processed as described previously (56), and the remaining supernatant was saved as the particulate fraction. Protein concentration was measured by BCA assay (Pierce Biotechnology). Protein extracts were run on 4%–12% Bis-Tris NuPage gels (Invitrogen), blooted onto nitrocellulose membranes, and probed with the following primary antibodies: rabbit polyclonal antibody generated against synthesized peptide KKPQITTEPHAT corresponding to RGS2 C terminus; Cn (diluted 1:500; BD Biosciences; ref. 21); Thr286–phospho-CaMKII (diluted 1:1,000; Affinity BioReagents); ERK, Thr202/Thr204–phospho-ERK; JNK, Thr183/Tyr185–phospho-JNK, p38, and Thr180/Tyr182–p38 (diluted 1:1,000; Cell Signaling Technology; ref. 21); PKG-1α (diluted 1:1,000); and GAPDH (diluted 1:3,000; IMGENEX or Cell Signaling Technology). Antibody binding was visualized by horseradish peroxidase–conjugated secondary antibodies and enhanced chemiluminescence (Pierce Biotechnology).

Quantitative real-time PCR. Total RNA was extracted from snap-frozen heart tissue using TRizol reagent (Invitrogen; ref. 21). The yield and purity of RNA was estimated spectrophotometrically using A260/A280 ratio. RNA (1 μg) was reverse transcribed into cDNA using SuperScript first-strand synthesis system (Invitrogen). cDNA was subjected to PCR amplification using TaqMan PCR Master Mix reagent (Applied Biosystems). TaqMan primers and probes for Rgs2, Nppa, Nppb, Myh7, Atp2a2, Rcan1, 18S, and Gapdh were purchased from Applied Biosystems.

Adult mouse myocyte preparation and immunohistochemistry. Mouse adult ventricular myocytes were isolated, fixed, and stained for confocal immunohistochemistry as described previously (57). Cells were plated on laminin-coated 6-well dishes in medium, and incubated in Tyrode’s solution with angiotensin (1 μM for 2 h; Sigma-Aldrich) or ET1 (0.5 μM for 2 h; Sigma-Aldrich) in the presence or absence of DT2 compound.
Protein synthesis of adult mouse cardiac myocytes. Protein synthesis was measured by [3H] leucine incorporation. After overnight culture, cells were stimulated with 100 nM ET1. After ET1 stimulation, [3H] leucine (1 μCi/ml) was added to the culture medium, and cells were further incubated for 24 h. The incorporated [3H] leucine was measured using a liquid-scintillation counter.

Statistics. All values are expressed as mean ± SEM. Group data were compared using 1- or 2-way ANOVA (with genotype and sham or TAC as categories) and Tukey’s post-hoc multiple-comparisons test for between-group differences. Comparisons between 2 groups were made using nonpaired 2-tailed Student’s t test. A P value less than 0.05 was considered significant. Sample sizes and individual statistical results for all analyses are provided in the figures, supplemental figures, and supplemental tables.

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