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Constitutively Activating GNAS Somatic Mutation in Right Ventricular Outflow Tract Tachycardia

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Abstract

Background: The cellular mechanism of focal, idiopathic right ventricular outflow tract (RVOT) tachycardia is thought to be due to cAMP-mediated triggered activity. A potential molecular mechanism has not yet been determined. We identified and characterized a novel missense somatic mutation in the gene (GNAS) encoding the stimulatory G protein ($G_s\alpha$) from a patient with RVOT tachycardia that is proposed to be the etiology of the clinical tachycardia.

Methods: Percutaneous endomyocardial biopsies were obtained from multiple sites in a patient with non-exertional, repetitive monomorphic RVOT tachycardia. Sequencing of extracted genomic DNA identified a $G_{s\alpha}$ W234R variant only at the site of tachycardia origin. Functional studies using *in vitro* transfection with S49 cyc⁻ murine lymphoma cells and measurement of cyclic AMP levels were performed. A trypsin protection assay assessed GTP binding kinetics and structural modeling predicted the impact of the mutation on protein-protein interactions. Whole cell patch clamp experiments of transfected CHO cells assessed the downstream effects of the mutation.

Results: *In vitro* studies of the GNAS mutation (W234R) demonstrated basal levels of cAMP ~16-fold higher than wild-type cells, consistent with constitutive stimulation of $G_s a$. Mutant $G_s a$ was partially protected from proteolysis after incubation with GTP, indicating diminished GTPase activity and reduced GTP hydrolysis as the mechanism for increased basal intracellular

Disclosures: None

Supplemental Materials: Supplemental Table I

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cAMP levels. Transfected mutant CHO cells increased unstimulated mean peak L-type calcium channel current density by ~50% and in silico modeling demonstrated spontaneous delayed afterdepolarizations and triggered activity.

Conclusions: We identified a novel somatic mutation in GNAS associated with RVOT tachycardia. The mutation results in constitutive activation of $G_s\alpha$, impairs GTP hydrolysis, and elevates basal cAMP levels, leading to enhanced L-type calcium current and triggered activity. These findings confirm that RVOT tachycardia can be caused by somatic mutations in signal transduction proteins that regulate intracellular cAMP and its downstream effectors.

Journal Subject Terms:

arrhythmias; electrophysiology; mechanisms; translational studies

Keywords

arrhythmia; GTP - binding proteins; mutation; signal transduction; ventricular tachycardia

Introduction

Approximately 10% of patients with ventricular tachycardia (VT) have no apparent structural abnormalities. The cellular and molecular mechanisms of ventricular tachycardia (VT) in the absence of structural heart disease, commonly referred to as idiopathic VT, are incompletely understood. Genomic mutations have been identified in ion channels and structural proteins that cause some of these ventricular arrhythmias, but the etiology often remains elusive for the most common form of idiopathic VT, a non-familial tachycardia that originates from a discrete focal site within the right ventricular outflow tract (RVOT).¹ Affected individuals present with either exercise-induced sustained VT and/or resting repetitive (nonsustained) monomorphic VT. The etiology of this arrhythmia has been thought in some patients to reflect a somatic mutation.^{2,3} An inability to consistently access the minute focus of myocardial tissue that causes these arrhythmias has impeded efforts to identify other potential somatic mutations.

Idiopathic RVOT tachycardia is sensitive to perturbations that lower intracellular calcium and stimulated levels of cAMP via a G protein-coupled mechanism. Therefore, the tachycardia terminates with adenosine (through inactivation of adenylyl cyclase via the A_1 adenosine receptor $[A_1AR]$ and $G_i\alpha$), with beta-blockers (via competitive inhibition of the beta-adrenergic receptor), or with edrophonium/vagal maneuvers (through activation of the M2 muscarinic receptor). RVOT tachycardia also terminates via direct blockade of the L-type calcium channel. These composite responses suggest that RVOT tachycardia is caused by cAMP-mediated intracellular calcium overload that gives rise to delayed afterdepolarizations and triggered activity.^{1,4–6} Based on these data and our previous findings of myocardial somatic mutations in the genes that encode the $G_i\alpha$ and the A_1AR in unique subtypes of RVOT tachycardia,^{2,3} as well as known constitutively active somatic mutations in GNAS that elevate basal levels cAMP in endocrine tumors,⁷ we hypothesized that cAMP-mediated RVOT tachycardia could be caused by a somatic mutation in GNAS.

Methods

The data that support the findings of this study are available from the corresponding author upon reasonable request.

Index Case

A 67-year-old man presented with repetitive monomorphic VT at rest. The longest spontaneous run of VT was 17 beats. Cardiac evaluation included a normal ECG and signal averaged ECG. Cardiac catheterization disclosed no significant coronary artery disease and left ventricular ejection fraction was 50%. There was no family history of VT or sudden cardiac death. During electrophysiologic study, sustained monomorphic VT was reproducibly induced with rapid ventricular pacing either in the presence or absence of concurrent isoproterenol infusion. The VT morphology was similar to the patient's spontaneous arrhythmia, having a left bundle branch block (LBBB), inferior axis configuration with a precordial transition in lead V_3 . VT terminated with rapid ventricular pacing, adenosine, verapamil and edrophonium. The site of origin of tachycardia was localized to the septal RVOT by endocardial catheter activation mapping. VT was successfully ablated by application of radiofrequency energy (30 W for 60 seconds).

Genetic analysis

Prior to ablation and after informed consent was obtained, percutaneous endomyocardial biopsies were obtained with a bioptome from multiple right ventricle sites as part of a Cornell University Medical College Institutional Review Board approved protocol. A single myocardial biopsy sample was obtained from the site of origin of arrhythmia in the RVOT (guided by multiplane fluoroscopy) and from 2 sites in the right ventricular apex (remote site). The biopsy specimens were frozen rapidly and stored for molecular analysis. Genomic DNA was prepared using proteinase K digestion and phenol-chloroform extraction.⁸ Using 25 ng of the patient's genomic DNA as a template, all exons corresponding to the coding region of $G_s \alpha$ and their flanking intronic sequences were PCR amplified [(94°C × 60s, $60^{\circ}\text{C} \times 60\text{s}, 72^{\circ}\text{C} \times 60\text{s}) \times 35$ cycles, $72^{\circ}\text{C} \times 7$ min] using $[\alpha - 3^{3}P]d\text{CTP}$ and rTth DNA Polymerase XL (Applied Biosystems). Novel and previously described⁹ primer sequences are listed in Supplemental Table 1. ³²P-labeled PCR products were denatured and studied by single strand conformation polymorphism analysis (SSCP) using 6% polyacrylamide gels in the presence or absence 5% (v/v) glycerol as previously described, 6,10 When a GNAS SSCP variant was identified, the PCR product containing the variant was subjected to PCR amplification and cloned into a sequencing vector. Approximately 10 clones from each sample were sequenced with bidirectional DNA sequence analysis using Big Dye terminators on an ABI 377 automated sequencer (Applied Biosystems).

Functional studies of the G_{as} W234R mutation

Intracellular cAMP Accumulation

Mutated GNAS, identified in genomic samples, was introduced into full length GNAS cDNA via site-directed mutagenesis kit (QuikChange XL, Stratagene) with oligonucleotides 5'-CGATGAACGCCGCAAGCGATCCAGTGCTTCAACG-3' and 5'-

CGTTGAAGCACTGGATCCGCTTGCGGCGTTCATCG-3[']. Mutant and wild-type cDNAs were subcloned into pCEP4 plasmid and transiently transfected into G_sa -deficient S49 cyc⁻ murine lymphoma cells (provided by Dr. Xin-Yun Huang, Cornell Medicine).¹¹ After 48 hours, S49 cyc⁻ expression of G_sa isoforms was analyzed by Western blot with anti- G_sa antibody (Santa Cruz Biotechnology). Bound antibody was detected via an ECL Advance Western Blotting Detection Kit (Amersham Biosciences). Parallel transfected cultures were treated with isoproterenol (100 μ M) and intracellular cAMP levels were assayed by a cAMP-¹²⁵I radioimmunoassay kit (DuPont NEN, Boston, MA).¹²

Expression and Purification of Recombinant Gsa in Sf9 Cells

Recombinant wild-type and mutant (W234R) $G_s \alpha$ proteins were produced using a baculovirus expression system. Full-length GNAS cDNA encoding wildtype or W234R was amplified by PCR and cloned into pFastBac donor plasmid (Invitrogen) along with an NH₂-terminal hexahistidine tag using the following oligonucleotides: 5'-ATGCATCACCATCACCATCACATGGGC-3' and 5'-TTAGAGCAGCTCGTACTGACGAAGGTG-3'. All inserts were verified by automated sequence analysis. Plasmids were transfected into DH10Bac cells to generate recombinant bacmid DNA per the instructions of the manufacturer (Invitrogen). $G_s \alpha$ baculovirus was harvested from Sf9 insect cells transfected with recombinant bacmid DNA and then used to infect additional Sf9 cells in order to prepare recombinant wild-type or W234R mutant $G_s \alpha$ protein. Recombinant proteins were purified from infected Sf9 cell lysates by Ni²⁺-NTA affinity chromatography (Invitrogen).¹³

Trypsin Protection Assay

Wild-type or mutant $G_s \alpha$ protein (1.0 mg/ml) was incubated for 60 minutes at room temperature with 100 μ M GTP or GTP γ S. Trypsin was added to a final concentration of 100 μ g/ml. After incubation for an additional 10 minutes at room temperature, the digested protein was analyzed by SDS-PAGE.¹⁴

GTP_γS Binding and GTP Hydrolysis

GTP binding (k_{app}) and hydrolysis (k_{cat}) were determined as previously described.¹⁴ Values for k_{cat} were calculated directly from a single round of GTP hydrolysis, performed in triplicate experiments, by allowing recombinant G-protein to bind GTP in the absence of Mg^{2+} . Catalysis was initiated by addition of Mg^{2+} and rebinding of radiolabeled substrate was prevented by addition of excess unlabeled guanine nucleotide.

Structural Modeling

The effects of the W234R mutation on $G_s \alpha$ structure-function were probed by *in silico* modeling. A model for the mutant $G_s \alpha$ protein was generated from the crystal structure of bovine $G_s \alpha^{15}$ (Protein Data Bank accession code 1AZT) by substituting arginine for tryptophan at the conserved amino acid residue 234 and subsequent energy minimization was determined using the GROMOS96 force field in DeepView.¹⁶ Structural figures were generated with PyMol software (http://www.pymol.org). Predictive protein-

protein interactions were assessed using Mutabind2 software (https://lilab.jysw.suda.edu.cn/research/mutabind2/).¹⁷

Whole cell patch clamp electrophysiology

We transiently transfected Chinese Hamster Ovary (CHO) cells (ATCC, Manassas, VA) with cDNA encoding human cardiac L-type Ca^{2+} channel subunits ($\alpha 1$, $\beta 2a$ and $\alpha 2-\delta$) and wild-type or W234R human GNAS, using Superfect transfection reagent and CMV promoter-based expression vectors. The mutation was made using the QuikChange Multi Site-Directed Mutagenesis Kit (Stratagene, San Diego, CA). cDNA encoding eGFP was co-transfected to facilitate identification of transfected cells using an IX50 inverted microscope equipped with fluorescence optics (Olympus, Cypress, CA).

CHO cell expression studies were conducted using 6 independent transfections per group, for 6 separate days of patch-clamp recording performed over several weeks. On each day, fresh transfection mixtures were prepared to avoid recurring pipetting errors. Wild-type recordings were matched with those of the mutant each day to minimize variation from day-to-day transfection efficiency. Plasmid sequences were identical except for the point mutation to avoid variation from different promoter efficiencies. Individual cells were chosen for recording based on visual inspection of robust GFP expression as a marker of efficient transfection.

After 24–48 hours, we quantified the effects of the $G_s \alpha$ W234R mutation on basal channel function using whole-cell patch clamp at room temperature (~17–20 °C), and standard intracellular and bath solution except that BaCl₂ (20 mM) was included in the bath solution as a surrogate charge carrier for CaCl₂. Recordings were performed using a MultiClamp 700A amplifier, a Digidata 1300 Analogue/Digital converter, and pClamp9 software (Molecular Devices, San Jose, CA). Cells were held at –80 mV and then stepped to membrane potentials between –60 and +60 mV at 10 mV intervals, and then peak current density was measured and plotted against voltage. Currents were corrected for cell membrane area, quantified by measuring cell capacitance.

In silico modeling of GNAS mutation

For mathematical modeling, we used a human ventricular cell model.¹⁸ The input for the human ventricular cell model was parameter values. Output was transmembrane potential as a function of time. We used the parameter values to simulate normal (non-mutant) action potentials. This is the standard setup for which the model was developed and validated against a range of experimental data, including action potentials, calcium transients and ionic currents. The G_sa mutation was modeled by increasing the value of the parameter controlling permeability of the L-type Ca²⁺ current as determined in our experimental study. The mutation was modeled as a 50% and 100% increase in the permeability of the L-type calcium current. The protocol to induce delayed after depolarizations (DADs) consisted of 2 Hz pacing for 30 s, followed by a pacing pause.¹⁹

The type of model used in this study (a dynamic model representing a complex biological system) inevitably contains several simplifications and assumptions. For one, as is standard for this type of model, it contains four separate intracellular Ca^{2+} compartments

that each have a uniform Ca^{2+} concentration, although Ca^{2+} almost certainly changes more continuously within the cell, especially within the bulk intracellular space. Other simplifications include omission of chloride currents and phosphorylation dynamics.

Statistical Methods

Continuous variables are presented as mean±SD. Based on normality of distribution, pairwise comparisons of continuous variables were made using Student's t-test. Comparisons of more than two continuous variables were performed using ANOVA. Assumptions of normality of distribution and common variance were verified. Post-hoc pairwise comparisons of means were performed using the Scheffe test. P<0.05 was considered statistically significant. Statistics were performed using SPSS version 24 (IBM, Chicago, IL).

Results

Molecular Genetic Analysis

Percutaneous endomyocardial biopsies from the site of VT origin and remote right ventricular sites were obtained. Genomic DNA was extracted and all 13 exons corresponding to the coding region of GNAS and their flanking intronic sequences were PCR amplified. A single variant SSCP pattern in the amplicon of GNAS exon 9 was detected in our patient's DNA sample obtained from the VT site of origin. The PCR product containing the SSCP variant was identified, and the original sample was PCR amplified and cloned into a sequencing vector. Bidirectional automated sequence analysis of the exon 9 amplicon revealed a T to C transition at nucleotide 700 that substitutes arginine for tryptophan at amino acid residue 234 (W234R) (Figure 1A). Residue W234 is conserved across several species, including mouse, rat, dog, cow and human, and across other human G proteins, including Gi, Gt and Gq (https://blast.ncbi.nlm.nih.gov) (Figure 1B). No other variants were observed in GNAS amplicons from the RVOT biopsy sample, and no sequence variants were found in any amplicon from the right ventricular apex biopsy samples.

Functional studies of the W234R mutation in Gsa.

To evaluate the functional effects of $G_s \alpha$ W234R, we generated GNAS cDNA corresponding to this mutant allele by site-directed mutagenesis of wild-type GNAS cDNA. Mutant and wild-type GNAS cDNA were transfected into $G_s \alpha$ deficient S49 cyc⁻ lymphoma cells and synthesis of both recombinant proteins was confirmed by Western blot (Figure 2). To determine the physiologic activity of these G proteins, we assessed isoproterenol-mediated changes in cAMP levels in S49 cyc⁻ lymphoma cells transfected with wild-type, W234R $G_s \alpha$ and equal ratios (1:1) of wild-type and mutant construct. Three separate experiments for each combination of the three constructs were carried out in triplicate.

S49 cyc⁻ lymphoma cells transfected with W234R mutant GNAS demonstrated markedly elevated basal levels of cAMP (79 \pm 12 pmol/10⁷ cells), which were 15.8-fold higher than that observed in basal wild-type GNAS transfected cells (5.0 \pm 0.7 pmol/10⁷ cells), P<0.001. For cells transfected with equal ratios (1:1) of wild-type and mutant constructs, the intracellular levels of cAMP were 12-fold greater than control, 60 \pm 8 pmol/10⁷ cells

To further delineate the mechanism by which $G_s \alpha$ W234R increases basal intracellular cAMP levels, the structural and functional properties of recombinant wild-type and W234R $G_s \alpha$ proteins were assessed with a trypsin protection assay. When bound to GDP, wild-type $G_s \alpha$ subunits are digested by trypsin into small fragments. Such extensive proteolysis occurs when wild-type $G_s \alpha$ subunits are incubated with GTP since the latter is hydrolyzed to GDP by $G_s \alpha$ GTPase activity. In contrast, GTP γ S is relatively resistant to hydrolysis, and thus, in the presence of GTP γ S, wild-type $G_s \alpha$ is protected against trypsin proteolysis after incubation with GTP (Figure 3A), findings consistent with diminished GTPase activity and increased basal intracellular cAMP levels.

In studies that directly assess guanine nucleotide binding and hydrolysis, we observed no difference between GTP γ S binding to wild-type or to W234R G_sa (K_d 0.045±0.006 versus K_d = 0.043±0.007, P=0.46) (Figure 3B). However, mutant G_sa exhibited reduced GTP hydrolysis. k_{cat} for wild-type G_sa was 5.64±1.0 min⁻¹ and for G_sa W234R was 1.56±0.7min⁻¹, P<0.001. These data suggest that G_sa W234R is trapped in the GTP bound active state, allowing G_sa W234R to increase basal intracellular cAMP levels in the absence of an agonist.

Models of Mutant G_sa Structure

The crystallographic structure of $G_s \alpha$ in a complex with GTP γ S has been previously reported and provides a model to determine the conformational consequences of $G_s \alpha$ mutations¹⁵ (Figure 4). GTP binding to $G_s \alpha$ produces conformational changes in three switch regions (I-III), that in turn determine the interactions of $G_s \alpha$ with the G protein coupled receptor, $\beta\delta$ dimers and its effector, adenylyl cyclase. GTP hydrolysis is thought to occur by deprotonation of an attacking water molecule and transfer of a proton to an oxygen of the γ -phosphate.¹⁶ This process is facilitated by two catalytic residues in $G_s \alpha$: R201 and Q227. Both sites have been associated with constitutively activating endocrine mutations.^{20–25}

W234 is located at the center of an evolutionarily conserved hydrophobic core that stabilizes the position of switch II including Q227. Mutation of W234 substitutes an arginine and introduces a polar side chain into this hydrophobic environment. As a result, the region is predicted to destabilize the orientation of Q227 for GTP hydrolysis. Based on modeling,¹⁷ the G_{bind} conferred by the W234R mutation would be 2.36 kcal/mol, which is consistent with a destabilizing and deleterious mutation ($G_{bind} > 1.5$ kcal/mol). In addition, the W234R mutation places the positively charged arginine next to K53, a conserved positively charged residue of the P-loop motif that binds the substrate phosphates, which would lead to

repulsion between the two sites. The net result of these effects is that GTP binding occurs in a conformation unsuitable for hydrolysis (Figure 4).

Whole cell patch clamp electrophysiology

We quantified whole-cell Ba²⁺ currents using whole-cell patch clamp in CHO cells coexpressing cardiac L-type Ca²⁺ channel subunits with wild-type or W234R mutant G_sa. Compared to wild-type G_sa, co-expression with W234R mutant G_sa increased mean peak L-type calcium channel current density by 48±15 % (P<0.05; n =24 – 27 cells per group) (Figure 5).

In silico model of increased calcium channel current density due to W234R Gsa mutation

Simulations with wild-type $G_s \alpha$ did not elicit DADs or triggered activity. Simulating the effects of W234R $G_s \alpha$ mutation by increasing L-type calcium current permeability by 50% resulted in prolonged action potential duration but no DADs. However, an increase of 100% in L-type calcium current permeability caused multiple DADs and triggered a spontaneous action potential following cessation of pacing, secondary to calcium-induced calcium release from the ryanodine receptor (Figure 6).

Discussion

We identified a constitutively activating somatic GNAS mutation from the arrhythmogenic site in the RVOT from a patient with repetitive monomorphic VT. W234R is a nonconservative charge-changing $G_s \alpha$ mutation, which impairs GTP hydrolysis and elevates basal cAMP levels. In addition, transfection studies demonstrated an increase in calcium current due to the W234R $G_s \alpha$ mutation and *in silico* studies were consistent with the presence of DADs and triggered beats. Along with our previous observation that somatic mutations in the genes encoding $G_{i2}\alpha$ and A_1AR can cause atypical forms of RVOT tachycardia,^{2,3} the present findings further strengthen the proposal that RVOT tachycardia can be linked to somatic mutations in signal transduction proteins that elevate cAMP, leading to DADs and triggered activity

Repetitive monomorphic VT is characterized by brief paroxysms of nonsustained VT at rest, occurring in the absence of structural heart disease.^{6,26} These arrhythmias may become sustained under conditions associated with enhanced catecholamine stimulation and/or programmed stimulation, as demonstrated in our patient. Mapping studies confirm a discrete, focal outflow tract source. The tachycardia has features consistent with calcium-dependent, cAMP-mediated VT, i.e., termination in response to adenosine, vagal maneuvers, and blockade of the slow-inward calcium current. Our patient's tachycardia demonstrated these characteristics, suggesting a pathogenic mechanism that is $G_s\alpha$ -mediated and dependent upon increased intracellular calcium.

Heterotrimeric G proteins, consisting of α , β , and γ subunits, play a central role in a variety of intracellular signal transduction systems (Figure 7).²⁷ G proteins are activated by binding of agonists (e.g., catecholamines) to G protein coupled receptors, resulting in release of GDP and binding of guanosine triphosphate (GTP). Binding of GTP results in conformational changes in three switch regions, causing the GTP- α complex to dissociate from both the

receptor and noncovalently bound β and γ subunits. The activated α subunit and the $\beta\gamma$ dimer regulate the activity of downstream effectors, such as AC, phospholipases and ion channels. This activity is terminated by intrinsic GTPase activity in the α subunit, which hydrolyzes GTP to GDP, reconstituting the inactive heterotrimeric complex. In the case of cardiac myocytes, the beta-adrenergic receptor is associated with the heterotrimeric G protein G_s α ; the M2 muscarinic receptor and A₁AR are associated with G_i α .

Functional studies of mutant W234R $G_s \alpha$ transfected into $G_s \alpha$ -deficient S49 cyc⁻ murine lymphoma cells showed a marked increase (approximately 16-fold) in basal levels of cAMP compared to cells transfected with wild-type $G_s \alpha$ (Figure 2). Moreover, basal levels of cAMP of mutant $G_s \alpha$ approximated adrenergically-stimulated (via isoproterenol) levels of cAMP in wild-type $G_s \alpha$ transfected cells. These findings suggest that the somatic mutation obtained from the site of VT origin caused constitutive activation of $G_s \alpha$. This was further supported by detection of reduced GTPase hydrolysis of mutant W234R $G_s \alpha$ (Figure 3). Furthermore, structural modeling predicted that the W234R mutation causes GTP binding in a conformation unsuitable for hydrolysis (Figure 4). Finally, whole cell patch clamp electrophysiology of mutant W234R $G_s \alpha$ increased L-type calcium channel current density by approximately 50% (Figure 5), and when further augmented, as would occur with adrenergic stimulation, DADs and triggered activity were simulated in an *in silico* model of human ventricular myocytes (Figure 6).

The findings above are consistent with the arrhythmic phenotype in our patient. The presenting arrhythmia, repetitive monomorphic VT, occurs at rest, but yet is cAMP-mediated.⁶ Agonist-independent G protein activation due to reduced GTPase activity may explain how cAMP- mediated tachycardia can occur in the absence of an apparent increase in endogenous or exogeneous catecholamine stimulation. Enhanced catecholamine stimulation, under appropriate conditions, may further augment elevated basal cAMP levels and facilitate initiation of sustained monomorphic VT. It is possible that compensatory mechanisms (i.e., enhanced phosphodiesterase activity or reduced adenylyl cyclase) may also be activated and prevent continuous VT despite persistent cAMP stimulation.

Although germline transmission of activating GNAS mutations is generally presumed to be lethal, somatic mutations, especially with restricted tissue expression and receptor coupling can lead to more limited manifestations. Constitutively activating mutations of $G_s \alpha$ (R²⁰¹ and Q²²⁷) have been described in acromegaly, thyroid adenomas, the McCune-Albright syndrome, and less commonly in ovarian, testicular stromal Leydig cell tumors and Cushing's syndrome.^{20–25} These somatic mutations have biochemical consequences similar to W234R and are characterized by loss of GTPase activity, impaired hydrolysis of GTP, sustained stimulation of adenylyl cyclase and persistent elevation of cAMP. Based on these data, we hypothesize that similar gain-of-function mutations in the stimulatory G protein $G_s \alpha$ may be responsible for some forms of cAMP mediated, adenosine-sensitive RVOT tachycardia.

The mutation identified in GNAS as well as previously reported arrhythmogenic somatic mutations in GNAI3 ($G_{i2}\alpha$) and ADORA1, identified by sequencing DNA from a 1–2 mm³ biopsy sample from the arrhythmogenic focus, is consistent with there being more than

one single cardiac cell affected by a somatic G protein mutation.^{2,3} However, it remains unknown whether these somatic mutations occur during embryonic heart development in a single cell that clonally expands or whether they possibly occur during postnatal life in multipotent cardiac cells that retain the capacity to proliferate in the adult heart.²⁸ The significance and prevalence of somatic gene mutations in individuals with RVOT tachycardia merits further investigation.

Other arrhythmogenic cardiac somatic mutations, although rare, have also been identified in the gene encoding connexin 40 (GJA5) in 3 patients with atrial fibrillation.²⁹ Additionally, somatic mutations in genes encoding the transcriptional factors GATA4, GATA5 and GATA6 are increasingly recognized as being associated with some forms of congenital heart disease, including ventricular septal defects and tetralogy of Fallot.³⁰

Study Limitations

We performed targeted sequencing of extracted genomic DNA of all exons corresponding to the coding region of GNAS and their flanking intronic sequences to identify the W234R variant in this study. Although we cannot rule out the presence of modifying polymorphisms that could have been implicated in our patient's RVOT tachycardia, in *vitro* functional studies (i.e., cAMP assay, trypsin protection assay, whole cell patch clamping), structural modeling, and *in silico* modeling all suggest that the identified mutation causes phenotypic changes consistent with of our patient's VT. Moreover, the mutation was only found at the site of arrhythmia origin and not at remote right ventricular biopsy sites, indicating the focal and somatic nature of the mutation. The generalizability of our findings to other patients with RVOT tachycardia is unknown since myocardial biopsies are rarely performed in these patients and the focal origin of the mutation makes exact sampling of the critical tissue often fortuitous.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

Sources of Funding:

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Nonstandard Abbreviations and Acronyms:

A ₁ AR	A ₁ adenosine receptor
cAMP	cyclic adenosine monophosphate
DAD	delayed afterdepolarization
Gsa	stimulatory G protein alpha-subunit
Gia	inhibitory G protein alpha subunit
GDP	guanosine diphosphate

GTP	guanosine triphosphate
PCR	polymerase chain reaction
RVOT	right ventricular outflow tract
SDS-PAGE	sodium dodecyl sulphate-polyacrylamide gel electrophoresis
SSCP	single strand conformation polymorphism analysis
VT	ventricular tachycardia

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What Is Known?

- Right ventricular outflow tract (RVOT) tachycardia occurs in the absence of structural heart disease and is due to a non-reentrant mechanism.
- This form of tachycardia is usually sustained and induced with catecholamine stimulation and burst pacing. However, a variant manifests as repetitive monomorphic ventricular tachycardia and occurs independently of catecholamine stimulation.

What the Study Adds?

- A cardiac somatic mutation in GNAS was identified from an endomyocardial biopsy obtained from the site of origin of the tachycardia. The mutation increases basal levels of cAMP by approximately 16-fold compared with wild-type GNAS. Further characterization of the mutation showed diminished GTPase activity and impaired GTP hydrolysis. Transfection experiments showed that the mutant protein increases basal inward calcium current by ~50% and in silico modeling demonstrated delayed afterdepolarizations and triggered activity.
- These findings highlight the role of somatic cardiac mutations in genes that regulate cAMP and its downstream effects on intracellular calcium as a contributing mechanism for some forms of RVOT tachycardia.

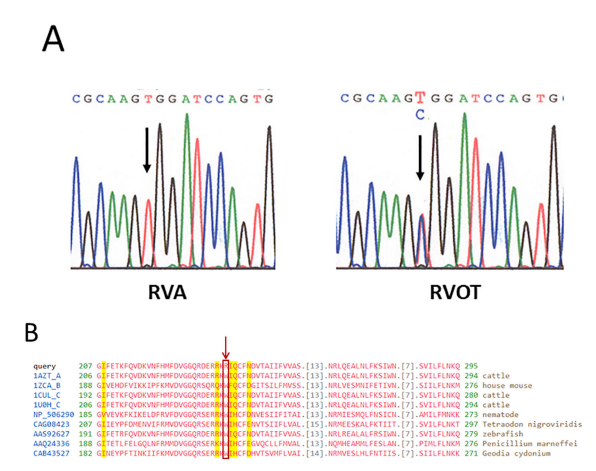


Figure 1.

Sequence of polymerase chain reaction-amplified DNA from right ventricular apex (RVA) and right ventricular outflow tract (RVOT) biopsy samples. **A**, Novel point mutation (W234R) in the switch II region of $G_s \alpha$ was identified from a biopsy sample obtained from the site of origin of ventricular tachycardia in the RVOT, but not from the RVA. **B**, $G_s \alpha$ protein sequence alignment showing site of point mutation (W234R) within a conserved region across several species.

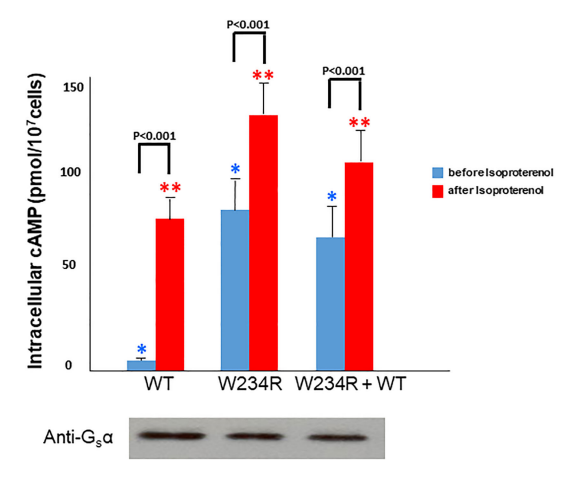


Figure 2.

Effects of W234R mutation in $G_s\alpha$ (stimulatory G protein alpha- subunit) on cyclic AMP (cAMP) cellular content. S49 cyc⁻ $G_s\alpha$ deficient cells were transfected with either wild-type or W234R mutant $G_s\alpha$ cDNA. Western blot demonstrated similar expression levels of both wild-type and mutant protein. cAMP levels were assayed before and after treatment with isoproterenol (ISO). Three separate experiments for each construct were carried out in triplicate. Thus, there are 9 data points for each condition. Basal cAMP levels in cells transfected with mutant $G_s\alpha$ were 5.0 ± 0.7 pmol/10⁷ cells, whereas basal cAMP levels in cells in cells transfected with mutant $G_s\alpha$ were 79 ± 12 pmol/10⁷ cells, P<0.001. * P< 0.001 for comparison of basal cAMP levels among the three constructs. ** P<0.001 for comparison of cAMP levels after treatment with ISO across all three constructs.



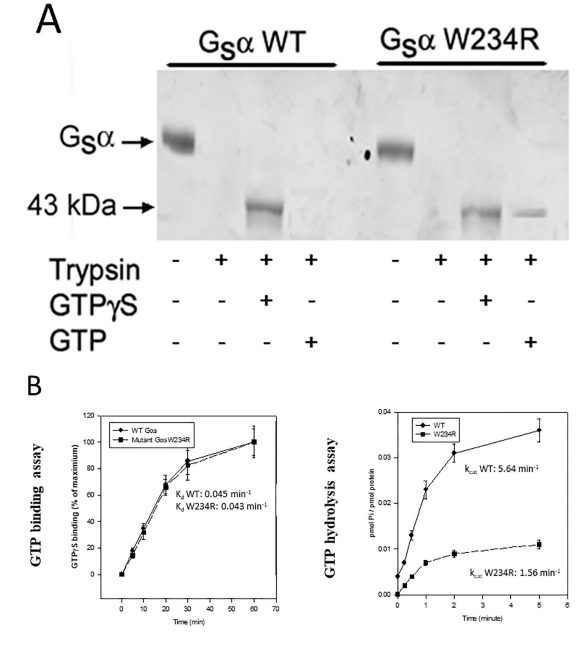


Figure 3.

Structural and functional properties of wild-type and mutant W234R $G_s \alpha$ recombinant proteins. **A**, Wild-type $G_s \alpha$ is protected from trypsin proteolysis after incubation with GTP γ S (43kDa proteolytic fragments accumulate) but not with GTP. **B**, Left panel. Recombinant wild-type and mutant $G_s \alpha$ proteins were incubated with ³⁵[S]GTP γ S and analyzed as described in Methods. No effect of mutation on GTP binding was observed and similar K_d values were calculated for both proteins (P=0.46). **B**, Right panel. Wild-type and mutant $G_s \alpha$ proteins were incubated with [γ -³²P]GTP. Aliquots (50 µl) were analyzed for [³²P]Pi at the indicted time. k_{cat} was calculated by observation of a single round of GTP hydrolysis performed in triplicate experiments.

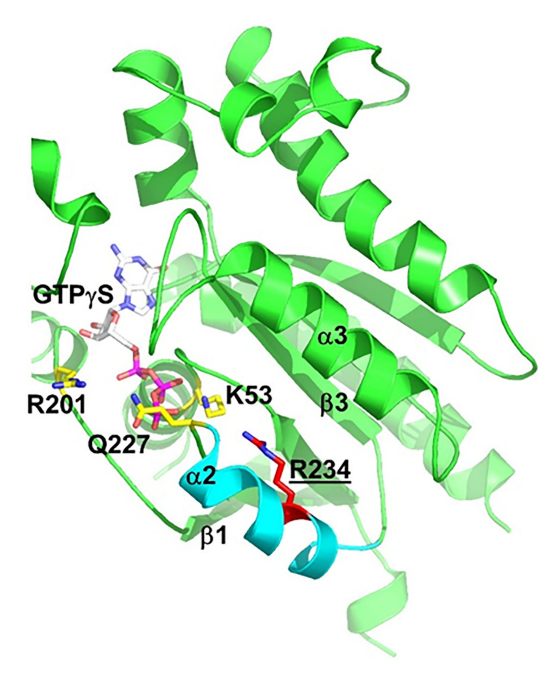


Figure 4.

Ribbon diagram for the Ras-like domain of bovine Gsa (PDB ID 1AZT), with W234 mutated to R (red sticks, label underlined). The bound GTP analog is colored according to atom types, and surrounding secondary structure elements are labeled according to Sunahara et al.¹⁵ Catalytically important residues are shown in yellow, and the switch II helix a2 in cyan.

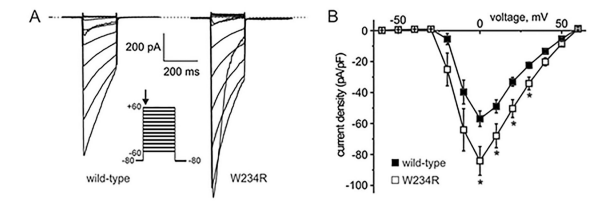


Figure 5.

Whole-cell patch clamp. **A**, Exemplar traces measured using whole-cell patch clamp from CHO cells expressing cardiac L-type calcium channel subunits ($\alpha 1$, $\beta 2a$ and $\alpha 2-\delta$) and wild-type or W234R human G_s α as indicated. Zero current level indicated by dashed line. Voltage protocol shown in lower inset. Arrow, approximate time point at which peak current was measured. **B**. Mean current density-voltage relationships for cells as in panel A; n = 27 (wild-type) and n = 24 (W234R). *P < 0.05 comparing wild-type versus W234R currents.

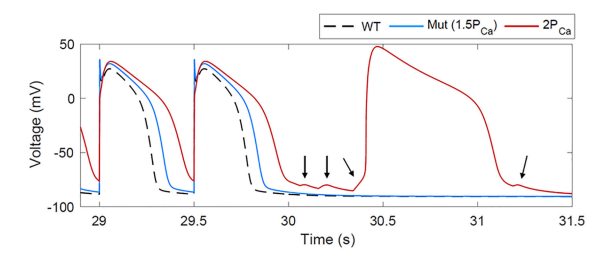


Figure 6.

Model prediction of proarrhythmic electrical activity due to increased L-type calcium current secondary to W234R $G_s \alpha$ mutation. Simulations of wild-type ionic behavior in response to 2 Hz pacing for 30 s followed by a pacing pause results in normal action potentials (black trace). Simulating the W234R $G_s \alpha$ mutation causes action potential prolongation (50% increase in the permeability of the L-type calcium current: blue trace). Challenging the cell by increasing the calcium current further leads to multiple DADs (arrows) and a spontaneously triggered action potential (100% increase in the permeability of the L-type calcium current; red trace).

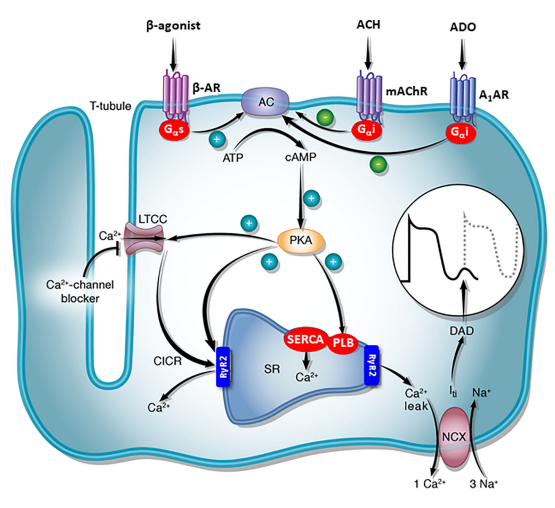


Figure 7.

Schematic showing the signal transduction cascade that mediates ventricular tachycardia due to cAMP-mediated triggered activity. Clinical right ventricular outflow tract tachycardia is usually adrenergically-mediated, and is frequently triggered by exercise or stress. In contrast, the somatic mutation W234R $G_s \alpha$ doesn't require an agonist for activation since it is constitutively active, accounting for resting ventricular tachycardia. Abbreviations: AC, adenylyl cyclase; A1AR, adenosine receptor A1; ACH; acetylcholine; ADO, adenosine; β -AR, β -adrenergic receptor; CICR, calcium-induced calcium release; DAD, delayed afterdepolarization; $G_{i2}\alpha$, inhibitory G-protein; $G_s\alpha$, stimulatory G-protein; I_{ti} , transient inward current; LTCC, L-type calcium channel; mAChR, muscarinic acetylcholine receptor; NCX, sodium–calcium exchanger; PKA, cAMP-dependent protein kinase (protein kinase A); PLB, phospholamban; RyR2, ryanodine receptor; SERCA, sarco/endoplamic reticulum Ca^{2+} -ATPase; SR, sarcoplasmic reticulum.