Gene Transfer of Engineered Calmodulin Alleviates Ventricular Arrhythmias in a Calsequestrin-Associated Mouse Model of Catecholaminergic Polymorphic Ventricular Tachycardia

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Background—Catecholaminergic polymorphic ventricular tachycardia (CPVT) is a familial arrhythmogenic syndrome characterized by sudden death. There are several genetic forms of CPVT associated with mutations in genes encoding the cardiac ryanodine receptor (RyR2) and its auxiliary proteins including calsequestrin (CASQ2) and calmodulin (CaM). It has been suggested that impairment of the ability of RyR2 to stay closed (ie, refractory) during diastole may be a common mechanism for these diseases. Here, we explore the possibility of engineering CaM variants that normalize abbreviated RyR2 refractoriness for subsequent viral-mediated delivery to alleviate arrhythmias in non–CaM-related CPVT.

Methods and Results—To that end, we have designed a CaM protein (GSH-M37Q; dubbed as therapeutic CaM or T-CaM) that exhibited a slowed N-terminal Ca dissociation rate and prolonged RyR2 refractoriness in permeabilized myocytes derived from CPVT mice carrying the CASQ2 mutation R33Q. This T-CaM was introduced to the heart of R33Q mice through recombinant adeno-associated viral vector serotype 9. Eight weeks postinfection, we performed confocal microscopy to assess Ca handling and recorded surface ECGs to assess susceptibility to arrhythmias in vivo. During catecholamine stimulation with isoproterenol, T-CaM reduced isoproterenol-promoted diastolic Ca waves in isolated CPVT cardiomyocytes. Importantly, T-CaM exposure abolished ventricular tachycardia in CPVT mice challenged with catecholamines.

Conclusions—Our results suggest that gene transfer of T-CaM by adeno-associated viral vector serotype 9 improves myocyte Ca handling and alleviates arrhythmias in a calsequestrin-associated CPVT model, thus supporting the potential of a CaM-based antiarrhythmic approach as a therapeutic avenue for genetically distinct forms of CPVT. (J Am Heart Assoc. 2018;7:e008155. DOI: 10.1161/JAHA.117.008155.)

Key Words: arrhythmia (mechanisms) • calcium channel • calcium signaling • calmodulin • gene therapy

Cardiac arrhythmia is a leading cause of mortality and morbidity worldwide.1,2 Aberrant Ca handling, and alterations in the cardiac sarcoplasmic reticulum (SR) Ca release channel (ryanodine receptor, RyR2), in particular, is recognized as an important factor in the genesis of arrhythmia.3,4 This link is especially evident in catecholaminergic polymorphic ventricular tachycardia (CPVT), an inherited arrhythmic syndrome caused by mutations in the RyR2 itself and its multiple accessory proteins including calsequestrin 2 (CASQ2) and calmodulin (CaM).5,6

Interestingly, previous work from our and other laboratories have demonstrated that impairment of the ability of RyR2

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Accompanying Figures S1 through S3 are available at http://jaha.ahajournals.org/content/7/10/e008155/DC1/embed/inline-supplementary-material-1.pdf

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**Clinical Perspective**

**What Is New?**
- Catecholaminergic polymorphic ventricular tachycardia is a life threatening inherited arrhythmia syndrome caused by mutations in the ryanodine receptor (RyR2) and its auxiliary proteins including calsequestrin 2 (CASQ2) and calmodulin (CaM).
- The underlying cause of catecholaminergic polymorphic ventricular tachycardia involves the compromised ability of RyR2 to stay closed during diastole thereby resulting in aberrant arrhythmogenic intracellular Ca release.
- Here we designed a CaM construct that combines enhanced Ca sensitivity with tighter RyR2 binding resulting in enhanced Ca-dependent inhibition of RyR2.

**What Are the Clinical Implications?**
- Gene transfer of this CaM construct (therapeutic CaM, TCaM) normalized myocyte Ca cycling and alleviated life-threatening arrhythmias in vivo in mice affected by a mutation within another disease-related protein (CASQ2).
- Thus, TCaMs may provide a therapeutic strategy for multiple genetic forms of catecholaminergic polymorphic ventricular tachycardia.

Methods

The data and analytic methods will be made available to other researchers for purposes of reproducing the results or replicating the procedure. The data that support the findings of this study are available from the corresponding author on reasonable request.

Ethical Approval

All animal procedures were approved by the Ohio State University Institutional Animal Care and Use Committee. The study conformed to the Guide for the Care and Use of Laboratory Animals published by the US National Institutes of Health (NIH Publication No. 85-23, revised 2011).

Mouse Models

CASQ2 R33Q and knock out (KO) mice (3–6 months old, males) in the C57BL/6 genetic background were utilized in this study. All animal procedures were approved by The Ohio State University Institutional Animal Care and Use Committee. The study conformed to the Guide for the Care and Use of Laboratory Animals published by the US National Institutes of Health (NIH Publication No. 85-23, revised 2011).

Virus Production and Injection

Adeno- associated viral vector serotype 9 (AAV9) vectors were produced by co-transfecting HEK293 cells with three plasmids as previously described, which were then expanded in 10% DMEM medium. Five days after transfection, the cells were

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harvested for downstream purification using PEG precipitation and an iodixanol gradient protocol. The 40% iodixanol gradient band was collected and subjected to further purification. The virus titer was obtained via TaqMan (LifeTechnologies) based quantification. AAV9 (100 µL, containing 1 x 10^{11} viral genomes) was injected into adult male mice (12–14 weeks old) through the intra-thoracic cavity. After being lightly anesthetized by ≈1% isoflurane, mice were kept in supine position to allow clear access to the chest area. A needle (29.5 gauge) was then inserted at an angle halfway between the ribs and ≈7.5 mm left of sternum. Care was taken to avoid insertion of the needle into the lungs or heart. This straightforward technique ensures efficient and highly reproducible viral transduction specifically to the adult heart, meanwhile minimizing the risk of damaging the beating heart.

### Measurement of the Rate of Ca Dissociation From CaM

Wild type (WT) and recombinant CaM were expressed in bacteria (DE3 BL21) and purified as previously described. Ca dissociation rates were measured using an Applied Photophysics Ltd. (Leatherhead, UK) model SX.18 MV stopped-flow instrument with a dead time of 1.4 ms at 20°C. Each rate represents an average of at least five traces.

CaM^{F20W} was utilized to record the rate of Ca dissociation from the N-terminal domain of isolated CaM, the samples were excited using a 150 W xenon arc source. The tryptophan (Trp) emission was monitored through a UV-transmitting black glass filter (UG1 from Oriel [Stanford, CT]). The data collected from CaM^{F20W} were fit with a single exponential. Each koff represents an average of at least five traces. The fluorescent Ca chelator Quin-2 was utilized to record the rate of Ca dissociation from CaM-binding domain in the presence of the human RyR2. CaM-binding domain (RSKKAVWHKLLSKQRKRAVCFM-RPLYNL). Data were fit with a double exponential (variance less than 1.3 x 10^{-4}) to account for the fact that Quin-2 reports the rates of Ca dissociation from both the N- and C-domains of CaM. The buffer used in all stopped-flow experiments was 10 mmol/L MOPS, 150 mmol/L KCl, pH 7.0.

### Cardiomyocyte Isolation and Confocal Ca Imaging

#### Myocyte isolation

Mice were fully anesthetized using 4% isoflurane in 95% oxygen, before surgically removing the heart. Mouse ventricular myocytes were isolated as previously described. Briefly, the hearts were quickly excised and perfused on a Langendorff’s apparatus at 37°C. After 5 minutes of perfusion with nominally Ca-free perfusion solution (containing, in mmol/L: 140 NaCl, 5.4 KCl, 0.5 MgCl_2, 10 Hepes, and 5.6 glucose [pH 7.3]), the perfusate was switched to perfusion solution containing Liberase TH (0.24 U; Roche) for the digestion of the connective tissue. After digestion, single ventricular myocytes were isolated from the dissected and triturated ventricles and stabilized in perfusion solution containing BSA (20 mg/mL).

#### Ca imaging in intact myocytes

The ventricular myocytes were loaded with 8 µmol/L Fluo-3 AM (Invitrogen, Carlsbad, CA) for 25 minutes at room temperature, followed by 25 minutes of incubation in fresh perfusion solution (de-esterification). Fluo-3 was excited with the 488 nm line of an argon laser and emission was collected at 500 to 600 nm. Fluo-3 fluorescence was recorded in the line-scan mode of the confocal microscope (Olympus Fluoview 1000). The myocytes were field-stimulated using extracellular platinum electrodes.

#### Patch-clamp recordings

For the voltage-clamp experiments the external solution contained in mmol/L: 140 NaCl, 5.4 CsCl, 2.0 CaCl_2, 0.5 MgCl_2, 10 HEPES, and 5.6 glucose (pH 7.4). Patch pipettes were filled with a solution that contained in mmol/L: 123 CsCl, 20 TEACl, 5 MgATP, 5 NaCl, 1 MgCl_2, 0.1 Tris GTP, 10 HEPES, and 0.1 Rhod-2 K-salt (Molecular Probes, OR) (pH 7.2). Rhod-2 dye was excited with 561 nm laser and fluorescence was collected at 570 to 620 nm wavelengths. Ca currents and corresponding Ca transients were evoked by 300 ms depolarizing steps from −50 to −40 ... 60 mV in 10 mV intervals. The excitation-contraction (EC) coupling gain was calculated as a ratio of Ca transient amplitude to the peak density of Ca current.

#### Ca imaging in permeabilized myocytes

The myocytes were permeabilized with saponin (0.01% for 50 seconds) dissolved in the internal solution, which contained (mmol/L): 120 potassium aspartate, 20 KCl, 0.81 MgCl_2, 1 KH_2PO_4, 0.1 EGTA, (free [Ca] 120 nmol/L), 20 µmol/L cAMP, 3 MgATP, 10 phosphocreatine, 20 Hepes (pH 7.2) and 5 U/mL creatine phosphokinase. The cells were incubated with either WT or mutant CaM for 25 minutes to allow equilibration of CaM binding to its targets. Ca sparks were detected and analyzed using a custom MATLAB (2014b,
The MathWorks, Inc, MA) script as described previously. Refractoriness factor (RF), calculated as the inverse of frequency of Ca sparks that occurred within 1 seconds following Ca wave, was used to characterize refractoriness of the SR Ca release.

Statistical Analysis
Results are expressed as Mean±SEM. Statistical significance was determined using either one-way ANOVA or unpaired Student t test. For certain data sets with smaller sample size, The Wilcoxon rank sum or Kruskal–Wallis test was applied. A P<0.05 was considered statistically significant.

Results
CPVT CaM Mutants Shorten RyR2 Refractoriness
Genetic causes of certain forms of CPVT (eg, mutations in the luminal RyR2 accessory protein CASQ2) involve premature reactivation of RyR2 via abbreviation of the time RyR2 remains refractory. Recently, mutations in CaM have also been shown to cause CPVT. Considering CaM binding to the cytosolic surface of RyR2 has been reported to contribute to inactivation of RyR2, we tested whether arrhythmogenic mutations of CaM also act by shortening RyR2 refractoriness. Ca imaging was performed in permeabilized cardiac myocytes (cytosolic Ca buffered at 120 nmol/L) supplemented with different CaM protein variants. Consistent with previous studies, the addition of the CPVT CaM N98S (100 nmol/L), to permeabilized myocytes isolated from WT mice significantly increased the frequency of Ca waves (as compared with WT CaM; Figure 1A and 1C). RyR2 refractoriness in permeabilized myocytes was assessed by examining sparks restitution following Ca waves. Refractoriness factor (RF) was calculated as inverse to spark frequency occurring within 1-second following Ca wave. RyR2 refractoriness was significantly shortened by CaM N98S compared with WT CaM (Figure 1C). Similar results were obtained with the CPVT CaM N54I (Figure S1). Together with previous results on CASQ2 and RyR2-mediated CPVT, these data suggests that shortened RyR2 refractoriness may be a common mechanism that promotes arrhythmias caused by both CASQ2 and CaM mutants.

To determine whether these two different regulators of RyR2 are additive or if they operate independently, we examined the impact of arrhythmogenic CaMs in myocytes derived from CPVT mice devoid of CASQ2 (CASQ2-KO). As expected, in the presence of WT CaM permeabilized CASQ2-KO myocytes displayed significantly higher frequency of Ca waves when compared with myocytes from WT animals (0.54±0.01 versus 0.37±0.02 per seconds, P<0.001). Notably, supplementation of permeabilized CPVT myocytes with CaM N98S further increased the frequency of Ca waves and shortened the refractory period in these cells (Figure 2). Thus, CPVT mutations in CaM and CASQ2 act in parallel to modulate RyR2 refractoriness. These results suggest that it may be possible to modulate the composite RyR2 refractoriness by targeting either one of these proteins. We examined the possibility of modulating RyR2 refractoriness through modifying the Ca-binding properties of CaM.
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Proteins is known to in protein (Figure 3B and 3C). Since association with target dissociation in vitro in WT and mutant recombinant CaM during diastole. To test this hypothesis we measured CaM Ca of RyR2 by CaM (Figure 3A).37

Structurally CaM is thought to be composed of a C-terminal domain that controls the docking of CaM to RyR2 and an N-terminal domain that governs the Ca-dependent regulation of RyR2 by CaM (Figure 3A).37-39 Based on this model, we hypothesized that altering (ie, accelerating or slowing) the release of Ca from the N-terminal domain of CaM should accordingly change CaM’s ability to help keep RyR2 refractory during diastole. To test this hypothesis we measured CaM Ca dissociation in vitro in WT and mutant recombinant CaM protein (Figure 3B and 3C). Since association with target proteins is known to influence CaM’s Ca binding kinetics,40 we performed experiments with the CaMs alone (Figure 3B) and CaMs complexed with a peptide corresponding to the CaM binding domain of RyR2 (CaM-pRyR2) (Figure 3C). In accordance with our prediction, the CPVT associated mutation N54I accelerated the rate of Ca dissociation of N-domain of isolated CaM (Figure 3B). Furthermore, the N54I mutation, in addition to shortening RyR2 refractoriness (Figure S1) accelerated the N-terminal rate of Ca dissociation from CaM when complexed with the RyR2 peptide (Figure 3C). In the presence of the target peptide, the values of the Ca dissociation rates (ranging from 10.6 to 19.0/second) were now comparable to Ca spark restitution rates in myocytes in the presence of the corresponding CaM variants (≈3/second).

To test further the notion that the rate of Ca dissociation from CaM’s N-domain contributes to CaM regulation of RyR2 refractoriness, a CaM construct (CaM D57A) with hindered ability to bind Ca at the N-terminal was engineered and its effect on SR Ca release was examined. The D to A point mutation in CaM at position 57 (CaM D57A) greatly accelerated the Ca dissociation from the N-terminal domain of CaM (Figure 4A). Similar to the CPVT CaMs, supplementation of CaM D57A to WT myocytes significantly increased the frequency of Ca waves, and shortened RyR2 refractoriness compared with WT CaM (Figure 4B and 4C). Thus, RyR2 refractoriness can be modulated by altering the N-terminal Ca dissociation rate of CaM. This property of CaM makes it an excellent therapeutic target for resetting CPVT associated abbreviated RyR2 refractoriness.

Engineered Therapeutic CaMs

Based on the above mentioned results, stabilizing Ca binding to CaM by slowing its rate of dissociation should prolong refractoriness and may potentially be therapeutic against arrhythmias. To slow CaM’s N-terminal rate of Ca dissociation, we stabilized its Ca binding induced open conformation by mutating a key hydrophobic residue, Met 37, into a polar residue Gln. Based on our previous studies on CaM’s evolutionary relative troponin C, a similar mutation in troponin C facilitates the Ca-induced structural transition of the protein, thus stabilizing Ca binding.42,43 As shown in Figure 5A, mutating Met 37 to Gln (CaM M37Q) drastically slowed the rate of N-terminal Ca dissociation from CaM. As predicted, addition of CaM M37Q to permeabilized CPVT myocytes (CASQ2 R33Q44) reduced Ca waves (Figure 5B and 5C), and prolonged the refractory period of RyR2 (Figure 5C) relative to WT CaM.

Previous reports have suggested that aberrant RyR2 Ca release in myocytes derived from failing hearts can be blunted by addition of a CaM with enhanced binding affinity to RyR2, GSH-CaM.45,46 Unexpectedly, the addition of GSH-CaM to the CPVT myocytes had no effect at preventing the adverse Ca waves and accordingly had no significant effect at prolonging the pathologically abbreviated refractory period in these myocytes (Figure 5B and 5C). However, the addition of GSH to the N-terminus of CaM M37Q to help target our engineered CaM to RyR2 markedly enhanced its potentially beneficial effects in the CASQ2 R33Q myocytes (Figure 5B and 5C). One could surmise that prolonging refractoriness with this GSH-M37Q-CaM (termed therapeutic CaM [TCAm]) in CPVT mice harboring the arrhythmogenic CASQ2 R33Q mutation would

Figure 2. CPVT CaM N98S increased Ca waves frequency and shortened RyR2 refractoriness in CASQ2 KO myocytes. A, representative line-scan images of SCWs in permeabilized CASQ2 KO myocytes exposed to either 100 nmol/L of wtCaM or 100 nmol/L of N98S CaM. Cytosolic Ca was clamped at ≈120 nmol/L with the slow Ca buffer EGTA. B, Average data for SCW frequency and refractoriness factor obtained from CASQ2 KO ventricular myocytes permeabilized with 100 nmol/L wtCaM and 100 nmol/L of N98S CaM, respectively, n=31 to 38 cells, *P<0.05 vs wtCaM. Ca indicates calcium; CaM, calmodulin; CASQ2, calsequestrin; CPVT, catecholaminergic polymorphic ventricular tachycardia; EGTA, ethylene glycol-bis(β-aminoethyl ether)-N,N,N′,N′-tetraacetic acid; KO, knock out; RyR2, ryanodine receptor 2; SCW, spontaneous Ca waves.
be anti-arrhythmic in vivo. This hypothesis can be tested in mice harboring the CPVT CASQ2 R33Q mutation.

**Viral-Mediated Gene Delivery of TCaM Alleviates Arrhythmia Burden in CPVT**

In order to incorporate TCaM into the murine heart, we generated an AAV9-TCaM viral construct which was injected transthoracically into CPVT R33Q mice. The construct also contained a sequence encoding mCherry for transfection verification. Figure 6A demonstrates that ~40% of the ventricular myocytes could be transduced with our viral construct based on mCherry fluorescence. Consistent with the CPVT phenotype, myocytes isolated from R33Q mice transfected with our control mCherry AAV9 virus displayed frequent diastolic Ca waves, when stimulated at 0.5 Hz in the

**Figure 3.** The effect of CPVT CaM N54I on the rate of Ca dissociation from N-domain of CaM. A, Ca bound N-domain of CaM binds RyR2 and keeps it refractory following CICR. Meanwhile, CASQ2 senses luminal (Ca) decrease and contributes to RyR2 refractoriness. B, the time course of the decrease in Trp fluorescence as Ca dissociated from CaM<sup>F20W</sup> constructs. The data traces have been staggered and normalized for clarity. C, the time course of the increase in Quin-2 fluorescence as Ca dissociated from CaM-pRyR2 complex. The data traces have been staggered and normalized for clarity. Ca indicates, calcium; CaM, calmodulin; CASQ2, calsequestrin; CICR, calcium induced calcium release; CPVT, catecholaminergic polymorphic ventricular tachycardia; RyR2, ryanodine receptor 2; Trp, tryptophan.
presence of 100 nmol/L isoproterenol, a β-adrenergic ago-
nist (Figure 6B). Notably, the frequency of these arrhythmo-
egenic events was markedly reduced (≈6-fold) in AAV9-TCaM
myocytes (Figure 6B and 6C). Additionally, RyR2 refractori-
ness was significantly prolonged in AAV9-TCaM myocytes
consistent with the results obtained in permeabilized
myocytes (Figure 6C). Therefore, TCaM alleviated cellular
arrhythmogenesis in intact myocytes isolated from AAV9
infected R33Q mice. Of note, TCaM did not alter Ca transient
amplitude, baseline cytosolic Ca (Figure 6D) or the extent of
shortening in the same myocytes (Figure S2). Moreover TCaM
had no significant impact on inward Ca currents, Ca transients
and EC coupling gain in patch clamped R33Q myocytes
(Figure S3). Thus, while inhibiting arrhythmogenic diastolic Ca
waves, TCaM did not significantly alter EC coupling, systolic
Ca release or contractile function in R33Q myocytes.

To test the ability of AAV9-TCaM to reduce arrhythmia
propensity in vivo, we performed surface ECG measurements
in anesthetized CASQ2 R33Q mice challenged with epineph-
rine (Epi)+caffeine (Caf).27 Following IP injection of Epi and Caf,
only ≈10% of WT control mice developed episodes of VT, but
nearly all untransfected CASQ2 R33Q mice exhibited CPVT
(Figure 7A and 7B). Additionally, 7 out of 8 (88%) of these
mice died following arrhythmia induction. Notably, the AAV9-
TCaM infected mice had markedly reduced incidence of VT
following epinephrine and caffeine challenge. Moreover, the
majority of the mice, 7 out of 8 (88%), survived following the
Epi/Caf injection (Figure 7C). Thus, our rationally engineered
therapeutic protein, TCaM, alleviated life threatening ventric-
ular arrhythmias in the CASQ2 R33Q mice in vivo.

Discussion

In this study, we provided evidence that rationally engineered
CaM can be delivered to CPVT mice to treat arrhythmias. In
particular, we demonstrated that therapeutic CaM proteins
may provide a therapeutic strategy for multiple genetic
forms of CPVT and potentially other forms of Ca-dependent
arrhythmias associated with altered RyR2 function.

Following a Ca transient Ca release channels become
inactivated and remain unresponsive for a certain time
interval termed mechanical/Ca refractory period.36,47 Refrac-
toriness is modulated by a number of different factors,
including luminal Ca, CASQ2,21,48–50 RyR2 phosphorylation
and oxidation states.51–53 Here we demonstrated that this
process is also influenced by CaM and cytosolic Ca (Figure 8).
Moreover, we showed that a CaM protein, ie, TCaM, designed
to enhance RyR2 refractoriness can normalize pathologically
abbreviated refractoriness to inhibit arrhythmogenesis in
CPVT.

The potential role of CaM in regulating RyR2 has long been
recognized.17–20 CaM is associated with RyR2 at its cytosolic
regulatory domain and inhibits RyR2 activity at elevated Ca.18
Recently several mutations of CaM have been linked to
cardiac arrhythmia and specifically to CPVT.31 However,
precisely how and through what molecular steps CaM
regulates RyR2 mediated SR Ca release in cardiomyocytes
and how these processes are altered to cause arrhythmia

Figure 4. Engineered CaM variant D57A exhibited a faster rate
of Ca dissociation and mimicked the effects of CPVT CaMs. A, the
time course of the decrease in Trp fluorescence as Ca dissociated
from CaMF20W constructs. The data traces have been staggered
and normalized for clarity. B, representative line-scan images of
SCWs in permeabilized WT myocytes exposed to 100 nmol/L
wtCaM or 100 nmol/L D57A CaM. Cytosolic Ca was clamped at
≈120 nmol/L with the slow Ca buffer EGTA. C, Average data for
SCW frequency and refractoriness factor obtained from WT
mouse ventricular myocytes permeabilized with wtCaM and D57A
CaM, respectively, n=40 cells, *P<0.05 vs wtCaM. Ca indicates
calcium; CaM, calmodulin; cAMP, cyclic adenosine monophos-
phate; EGTA, ethylene glycol-bis[β-aminoethyl ether]-N,N,N',
N'-tetraacetic acid; SCW, spontaneous Ca waves; TCaM, thera-
peutic calmodulin; Trp, tryptophan; WT, wild type.

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remains to be elucidated. Here, we demonstrated that CaM contributes to RyR2 refractoriness through binding of Ca to CaM’s N-terminal domain and subsequent inhibition of RyR2 following SR Ca release. In support of this conclusion engineered mutations that accelerated Ca dissociation from the N-terminal domain of CaM shortened RyR2 refractoriness whereas engineered mutations that slowed Ca dissociation prolonged RyR2 refractoriness in permeabilized myocytes (Figures 4 and 5). Moreover, we also demonstrated that impairment of this regulatory mechanism accounts for arrhythmogenesis in CaM-dependent CPVT. In particular, our experiments showed that CPVT mutations in CaM accelerated Ca dissociation from the N-terminal domain of CaM and shortened RyR2 refractoriness (Figures 1 and 3; Figure S1). These findings are supported also by previous results of Sondergaard et al who showed that CPVT CaM N54I increased the rate of Ca dissociation from N-domain of CaM.

We and others have previously demonstrated that the generation of arrhythmogenic Ca release in CPVT involves shortened refractoriness of RyR2 in CPVT models associated with mutations in CASQ2.7,21 CPVT mutations in CaM and CASQ2 produced additive effects on RyR2 refractoriness and cellular arrhythmogenesis, supporting the notion that shortened refractoriness is a common mechanistic step in CPVT (Figure 2). Given the general role of RyR2 refractoriness in arrhythmogenesis, we reasoned that normalizing shortened RyR2 refractoriness by a rationally designed CaM with enhanced ability to induce RyR2 refractoriness may provide a therapeutic strategy for genetically distinct forms of CPVT. We were able to test this hypothesis by designing a therapeutic CaM that possessed a slowed N-terminal domain Ca dissociation rate and whose binding to RyR2 was enhanced by the GSH tag attached to the N-terminus of the protein. Introduction of this TCaM into permeabilized myocytes derived from mice affected by CASQ2-dependent CPVT normalized shortened refractoriness and reduced the frequency of diastolic Ca waves in these cells. Moreover viral-mediated transthoracic delivery of TCaM to adult CASQ2 R33Q mice reduced myocyte arrhythmic potential, in vivo susceptibility to arrhythmias and death in these mice.

Figure 5. TCaM significantly reduced spontaneous Ca waves in permeabilized R33Q myocytes and prolonged refractoriness. A, the time course of the decrease in Trp fluorescence as Ca dissociated from CaM$^{20W}$ constructs. The data traces have been staggered and normalized for clarity. B, representative line-scan images of SCWs in permeabilized R33Q myocytes exposed to cAMP. Cytosolic Ca was clamped at $\approx 120$ nmol/L with the slow Ca buffer EGTA. C, Average data for SCW frequency (n=18–39 cells) and refractoriness factor (n=11–15 cells) obtained from R33Q ventricular myocytes permeabilized with wtCaM, GSH-CaM, M37Q CaM, and GSH-M37Q CaM (TCaM), *P<0.05 vs wtCaM. Ca indicates calcium; CaM, calmodulin; cAMP, cyclic adenosine monophosphate; EGTA, ethylene glycol-bis(β-aminoethyl ether)-N,N,N′,N′-tetraacetic acid; SCW, spontaneous Ca waves; TCaM, therapeutic calmodulin; Trp, tryptophan; WT, wild type.

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Recent studies from Priori and coworkers demonstrated that wild type CASQ2 gene replacement utilizing AAV can rescue a CPVT phenotype associated with a mutated or absent CASQ2. However, such wild type replacement-gene transfer has the limitation that it can tackle only the subset of CPVT disorders dependent on aberrant CASQ2. Here we demonstrated that an engineered gene transfer of the RyR2 regulatory protein CaM can counter a non-CaM modulated arrhythmia model, suggesting that this therapeutic TCaM approach may serve as a general therapeutic avenue for CPVT mutations that result in aberrant Ca release. We have previously shown that abnormally shortened RyR2 refractoriness contributes to arrhythmogenesis associated with altered posttranslational modifications of RyR2 in settings of acquired cardiac disease, including post-infarction VF and non-ischemic heart failure. TCaMs could potentially serve as a general therapeutic approach to alleviate arrhythmia caused by various genetic and acquired molecular defects in the RyR2 channel complex.

It should be noted that TCaM proved highly effective at preventing arrhythmia despite a moderate transfection efficacy (≈40% myocytes transfected). Priori et al noted a similar unexpectedly high antiarrhythmic potential in the face of low transfection efficacy of virally transferred WT CASQ2. The unexpectedly strong antiarrhythmic effects of this delivery method could be explained by considering the role of synchronization of aberrant diastolic SR Ca release in the genesis of Ca-dependent arrhythmia. In order to initiate triggered activity in the form of triggered action potentials, aberrant Ca release has to occur synchronously across a critical mass of myocardium. We recently demonstrated that diastolic Ca release is indeed highly synchronized in CPVT ventricular myocytes and in cardiac tissue, due in part to

Figure 6. TCaM infected R33Q myocytes had improved Ca handling. A, evaluation of the infection efficiency of TCaM: phase contrast and mCherry fluorescence of TCaM injected R33Q myocytes. B, representative line-scan images of SCWs in intact myocytes isolated from mCherry or TCaM virus infected R33Q mouse. Myocytes were field-stimulated at 0.3 Hz in the presence of 1 μmol/L isoproterenol. C, Average data for SCW frequency (n=45–50 cells) and refractoriness factor (n=15–17 cells) obtained from R33Q myocytes infected with control (mCherry) and TCaM virus, respectively, *P<0.05 vs mCherry. D, Average of Ca transient amplitude and baseline fluorescence obtained from R33Q myocytes infected with control (mCherry) and TCaM virus, respectively, *P<0.05 vs mCherry. Ca indicates calcium; TCaM, therapeutic calmodulin; SCW, spontaneous Ca waves.

Figure 7. TCaM alleviated ventricular tachycardia in vivo in CPVT mice R33Q. A, Representative surface ECG traces from WT mice, R33Q mice or R33Q mice treated with TCaM. B, Average of VT incidence, n=8 to 19, *P<0.05 vs R33Q, #P<0.05 vs WT. C, Percentage of survival after ECG n=8 to 19, *P<0.05 vs R33Q, #P<0.05 vs WT. CPVT indicates catecholaminergic polymorphic ventricular tachycardia; ECG, electrocardiogram; TCaM, therapeutic calmodulin; VT, ventricular tachycardia.

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Figure 8. Multiple influences converge to impair RyR2 refractoriness and contribute to ryanopathies. CAM indicates calmodulin; CASQ2, calsequestrin; RyR2, ryanodine receptor 2.

Abnormally abbreviated Ca release refractoriness. Therefore, slowing Ca release refractoriness with TCaMs is expected to reduce the incidence of synchronized aberrant Ca release events on the cellular level, thus reducing triggered activity in tissue and arrhythmias in vivo.

In summary, this study demonstrates that a smartly formulated CaM variant designed to prolong RyR2 refractoriness inhibits arrhythmogenic aberrant Ca release in myocytes derived from CPVT hearts. Importantly, adenoviral-mediated thoracic cavity delivery of this CaM alleviated CPVT episodes and death in vivo. Our study points to a new therapeutic strategy for CPVT and possibly other forms of Ca-dependent arrhythmias using rationally engineered CaMs. Although we are discovering common aberrant mechanisms for arrhythmias, not all diseased hearts have the same etiology, genetic background or co-morbidities. Designer proteins open the doors for unprecedented personalized, and potentially, even generalized medicines as gene therapy or protein delivery techniques come to fruition.

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Disclosures
None.

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**Figure S1.** CPVT CaM N54I increased Ca waves frequency and shortened RyR2 refractoriness. 

A) Representative line-scan images of SCWs in permeabilized WT myocytes exposed to cAMP. 

B) Average frequency of SCWs (n=46-51 cells) and refractoriness (n=21-22 cells), *p<0.05 vs wtCaM. CPVT, Catecholaminergic polymorphic ventricular tachycardia; CaM, Calmodulin; Ca, Calcium; RyR2, Ryanodine receptor 2; cAMP, Cyclic adenosine monophosphate; SCW, spontaneous Ca waves; WT, wild type.
Figure S2. TCaM did not alter the extent of myocyte shortening in R33Q myocytes. **A**) representative shortening traces of R33Q myocytes infected with control (mCherry) or TCaM virus. **B**) Average percentage of shortening (n=45-47 cells), *p<0.05 vs mCherry. TCaM, Therapeutic Calmodulin.
Figure S3. EC-coupling in R33Q myocytes noninfected (control) and infected with TCaM. A) Voltage-dependence of Ca currents (I_{Ca}) and corresponding Ca transients recorded in control R33Q myocytes (n=6) and in R33Q myocyte expressing TCaM (n=4). B) Representative traces of Ca transients and I_{Ca} evoked by depolarizing steps from -50 to -20, 0, and 20 mV in control and TCaM myocytes, respectively. C) The EC-coupling gain in R33Q control and TCaM myocytes. *, P<0.05 vs control. EC, excitation contraction; TCaM, Therapeutic Calmodulin; Ca, Calcium; I_{Ca}, Calcium current.
Gene Transfer of Engineered Calmodulin Alleviates Ventricular Arrhythmias in a Calsequestrin–Associated Mouse Model of Catecholaminergic Polymorphic Ventricular Tachycardia

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