Ryanodine receptor–bound calmodulin is essential to protect against catecholaminergic polymorphic ventricular tachycardia

Yoshihide Nakamura,1 Takeshi Yamamoto,2 Shigeki Kobayashi,1 Masaki Tamitani,1 Yorimichi Hamada,1 Go Fukui,1 Xiaojuan Xu,3 Shigehiko Nishimura,1 Takayoshi Kato,1 Hitoshi Uchinoumi,1 Tetsuro Oda,1 Shinichi Okuda,1 and Masafumi Yano1

1Department of Medicine and Clinical Science, Division of Cardiology, and 2Faculty of Health Sciences, Yamaguchi University Graduate School of Medicine, Yamaguchi, Japan. 3Department of Pathology and Pathophysiology, School of Medicine, Tongji University, Shanghai, China.

Catecholaminergic polymorphic ventricular tachycardia (CPVT) is caused by a single point mutation in the cardiac type 2 ryanodine receptor (RyR2). Using a knockin (KI) mouse model (R2474S/+), we previously reported that a single point mutation within the RyR2 sensitizes the channel to agonists, primarily mediated by defective interdomain interaction within the RyR2 and subsequent dissociation of calmodulin (CaM) from the RyR2. Here, we examined whether CPVT can be genetically rescued by enhancing the binding affinity of CaM to the RyR2. We first determined whether there is a possible amino acid substitution within the CaM-binding domain in the RyR2 (3584–3603 residues) that can enhance its binding affinity to CaM and found that V3599K substitution showed the highest binding affinity of CaM to the CaM-binding domain. Hence, we generated a heterozygous KI mouse model (V3599K/+) with a single amino acid substitution in the CaM-binding domain of the RyR2 and crossbred it with the heterozygous CPVT-associated R2474S/+-KI mouse to obtain a double-heterozygous R2474S/V3599K-KI mouse model. The CPVT phenotypes — bidirectional or polymorphic ventricular tachycardia, spontaneous Ca2+ transients, and Ca2+ sparks — were all inhibited in the R2474S/V3599K mice. Thus, enhancement of the CaM-binding affinity of the RyR2 is essential to prevent CPVT-associated arrhythmogenesis.

Introduction

The cardiac type 2 ryanodine receptor (RyR2) is a huge tetrameric protein in the sarcoplasmic reticulum (SR) through which a large amount of Ca2+ is released from the SR, triggered by a small amount of Ca2+ entering the L-type Ca2+ channel (1, 2). To date, more than 150 mutations within the RyR2 have been reported to be linked with catecholaminergic polymorphic ventricular tachycardia (CPVT) (3). The point mutations observed in CPVT are not randomly distributed but clustered into 3 hot domains: the N-terminal (1–600 residues), the central (2000–2500 residues), and the C-terminal (4). Based on the “domain switch hypothesis” Ikemoto and colleagues proposed (5), we demonstrated that the single point mutation causes unzipping between the N-terminal and the central domains, which gave rise to a Ca2+ leak in the CPVT-associated R2474S/+-KI mouse (6) and also in a canine model of tachycardia-induced heart failure (7–9). These findings suggest that domain unzipping between the N-terminal and the central domains plays a critical role in the pathogenesis of lethal arrhythmia and heart failure as well. Recently, we found that domain unzipping displaces calmodulin (CaM) from the RyR2, which otherwise plays an inhibitory role in channel opening, causing Ca2+ leak in CPVT (10) and heart failure (11). Moreover, we found that Gly-Ser-His CaM (GSH-CaM), which has a higher binding affinity to the RyR2 than WT CaM, inhibited spontaneous Ca2+ sparks or transients as an arrhythmogenic substrate of inducing CPVT (12) and ameliorated contractile dysfunction in heart failure (13). These notions support the idea that the specific binding of CaM to the RyR2 plays a critical role in the pathogenesis of CPVT and heart failure. Furthermore, the fact that genetic modulation of the binding site of CaM to the RyR2 (3 amino acid substitutions: RyR2-
W3587A, -L3591D, -F3603A), which prevented CaM binding to the RyR2, induced premature death of the KI mice within 1 month, associated with marked cardiac enlargement and hypertrophy (14), also strongly indicates the essential role of CaM bound to the RyR2 in maintenance of cardiac function and structure. However, it remains to be elucidated as to whether forced association of CaM to the defective RyR2 rescues the diseased phenotypes. We hypothesized that the enhancement of CaM-binding affinity toward the defective RyR2 under stressful conditions prevents spontaneous Ca\(^{2+}\) release, thereby preventing CPVT.

### Results

A single amino acid substitution (V3599K) within the CaM-binding domain in the RyR2 (3584–3603 residues) markedly enhanced the binding affinity to CaM. We first determined whether there is a possible amino acid substitution within the CaM-binding domain in the RyR2 (3584–3603 residues) that can enhance its binding affinity. For this purpose, we assessed the binding of CaM to the RyR2 using a photoreactive cross-linker, sulfosuccinimidyl-6-(4′-azido-2′-nitrophenylamino)hexanoate (sulfo-SANPAH) (see Methods) to evaluate the CaM-binding affinity of 20-mer CaM-binding peptides (CaMBPs) harboring the same amino acid sequence as the CaM-binding domain (3584–3603 residues) except 1 amino acid residue (Table 1). Of the 25 peptides, peptide 24, in which a V3599K substitution was done, showed the highest inhibition of CaM-SANPAH binding to the RyR2, indicating a higher binding affinity of CaM to the 24 CaMBP than to the RyR2 (Figure 1). The conformational state change of CaM, upon its binding to CaMBPs, was also evaluated by measuring the tryptophan fluorescence change. As shown in Supplemental Figure 1 (supplemental material available online with this article; https://doi.org/10.1172/jci.insight.126112DS1), the V3599K CaMBP showed the highest change in CaM-mediated tryptophan fluorescence. The effect of a single amino acid substitution (V3599K) within the CaMBP on the binding affinity of CaM was also confirmed by quartz crystal microbalance (QCM) method. The single amino acid substitution in CaMBP (CaMBP mutation [CaMBPmut]: V3599K) markedly enhanced the binding of CaM to CaMBPmut. Suramin, which directly binds to the core RyR-CaM–binding sequence and displaces CaM from both RyR1 and RyR2 channel isoforms (15–17), indeed inhibited CaM binding to both WT CaMBP and CaMBPmut (Figure 2).

### Table 1. Single amino acid substitution within the CaM-binding domain in the RyR2

<table>
<thead>
<tr>
<th>Name</th>
<th>Amino acid sequence</th>
</tr>
</thead>
<tbody>
<tr>
<td>WT</td>
<td>AVWHKLQSKQRKRAVVACFR</td>
</tr>
<tr>
<td>1</td>
<td>AVWHRLQSKQRKRAVVACFR</td>
</tr>
<tr>
<td>2</td>
<td>AVWHYLQSKQRKRAVVACFR</td>
</tr>
<tr>
<td>3</td>
<td>AVWHKLQSKQRKRAVVACFR</td>
</tr>
<tr>
<td>4</td>
<td>AVWHKLQSKQRKRAVVACFR</td>
</tr>
<tr>
<td>5</td>
<td>AVWHKLQSKQRKRAVVACFR</td>
</tr>
<tr>
<td>6</td>
<td>AVWHKLQSKQRKRAVVACFR</td>
</tr>
<tr>
<td>7</td>
<td>AVWHKLQSKQRKRAVVACFR</td>
</tr>
<tr>
<td>8</td>
<td>AVWHKLQSKQRKRAVVACFR</td>
</tr>
<tr>
<td>9</td>
<td>AVWHKLQSKQRKRAVVACFR</td>
</tr>
<tr>
<td>10</td>
<td>AVWHKLQSKQRKRAVVACFR</td>
</tr>
<tr>
<td>11</td>
<td>AVWHKLQSKQRKRAVVACFR</td>
</tr>
<tr>
<td>12</td>
<td>AVWHKLQSKQRKRAVVACFR</td>
</tr>
<tr>
<td>13</td>
<td>AVWHKLQSKQRKRAVVACFR</td>
</tr>
<tr>
<td>14</td>
<td>AVWHKLQSKQRKRAVVACFR</td>
</tr>
<tr>
<td>15</td>
<td>AVWHKLQSKQRKRAVVACFR</td>
</tr>
<tr>
<td>16</td>
<td>AVWHKLQSKQRKRAVVACFR</td>
</tr>
<tr>
<td>17</td>
<td>AVWHKLQSKQRKRAVVACFR</td>
</tr>
<tr>
<td>18</td>
<td>AVWHKLQSKQRKRAVVACFR</td>
</tr>
<tr>
<td>19</td>
<td>AVWHKLQSKQRKRAVVACFR</td>
</tr>
<tr>
<td>20</td>
<td>AVWHKLQSKQRKRAVVACFR</td>
</tr>
<tr>
<td>21</td>
<td>AVWHKLQSKQRKRAVVACFR</td>
</tr>
<tr>
<td>22</td>
<td>AVWHKLQSKQRKRAVVACFR</td>
</tr>
<tr>
<td>23</td>
<td>AVWHKLQSKQRKRAVVACFR</td>
</tr>
<tr>
<td>24</td>
<td>AVWHKLQSKQRKRAVVACFR</td>
</tr>
<tr>
<td>25</td>
<td>AVWHKLQSKQRKRAVVACFR</td>
</tr>
</tbody>
</table>

The underlined letters represent the substituted amino acids.
Ventricular tachycardia was not induced by catecholamine or exercise in a R2474S/V3599K-KI mouse model.

We then generated a heterozygous V3599K/wt-KI mouse model in which the binding affinity of CaM to the RyR2 was markedly enhanced by a single amino acid mutation (V3599K) at the CaM-binding domain within the RyR2 (Supplemental Figure 2), and then we crossbred the V3599K/wt-KI mouse with the R2474S/wt-KI mouse to obtain a heterozygous R2474S/V3599K-KI mouse model. There was no appreciable difference in the structural or functional features of the hearts among WT (+/+), V3599K/wt-KI, and R2474S/V3599K-KI mice (structural features, Supplemental Figure 3; echocardiography, Supplemental Figure 4; protein expression and phosphorylation, Supplemental Figure 5).

We previously demonstrated that in the pathogenesis of CPVT, catecholamine-mediated phosphorylation of the RyR2 at serine 2808 (Ser2808) caused domain unzipping between the N-terminal and the central domains, which in turn dissociated CaM from the RyR2, inducing Ca\(^{2+}\) leak, leading to delayed afterdepolarization (DaD) and finally lethal arrhythmia (6, 10). In this study, we examined whether arrhythmia is observed in R2474S/V3599K-KI mice using 24-hour electrocardiography (ECG) monitoring and examined the effect of epinephrine or exercise on ventricular arrhythmia. Ventricular tachycardia (VT) was easily induced by epinephrine or exercise in all R2474S/+ mice but not in R2474S/V3599K-KI mice.
as well as WT and V3599K/+-KI mice (Figure 3, A and B). Under 24-hour ECG monitoring, both VT and ventricular bigeminy were frequently observed even under a small amount of stress in R2474S/+ mice but not mainly in R2474S/V3599K-KI mice (Figure 3, C and D).

Spontaneous Ca\(^{2+}\) transients and Ca\(^{2+}\) sparks were both inhibited in the R2474S/V3599K mice. There was no statistically significant difference in the contour or kinetic parameters of cell shortening (Figure 4) and Ca\(^{2+}\) transients (Figure 5A) at the baseline between the WT and V3599K/+-KI mice. In response to ISO, the time from the peak to an 80% decline in cell shortening or Ca\(^{2+}\) transients was prolonged in R2474S/+ mouse cardiomyocytes, suggesting a delay in the inactivation of Ca\(^{2+}\) or spontaneous Ca\(^{2+}\) release events. However, the time from the peak to an 80% decline in cell shortening or Ca\(^{2+}\) transients was virtually restored in R2474S/V3599K-KI cardiomyocytes (Figure 4 and Figure 5A). In the presence of ISO, the SR Ca\(^{2+}\) content increased in WT and V3599K/+-KI mice, while the extent of increase decreased in R2474S/+ mice, presumably because of Ca\(^{2+}\) leak (Figure 5B). On the other hand, the decrease in SR Ca\(^{2+}\) content in R2474S/+-KI mice was not observed in R2474S/V3599K-KI mice (Figure 5B). In the presence of ISO, the Ca\(^{2+}\) spark frequency increased in R2474S/+-KI mice compared with the R2474S/V3599K-KI mice (Figure 6A). Compared with the WT cardiomyocytes, both the peak amplitude and full width at half maximum (FWHM) decreased whereas full duration at half maximum (FDHM) increased in R2474S/+-KI cardiomyocytes, suggesting a delay in the inactivation of the RyR2. In contrast, the increase in FDHM was attenuated in R2474S/V3599K-KI cardiomyocytes (Figure 6B and Table 2). In the presence of ISO, spontaneous Ca\(^{2+}\) transients (SCaTs) increased in R2474S/+-KI mice, while it was hardly observed in R2474S/V3599K-KI mice (Figure 7). These findings clearly indicate that the defective interaction of CaM with the RyR2 and the subsequent Ca\(^{2+}\) leak causes not only an arrhythmogenic substrate leading to CPVT, but also of particular interest, a relaxation disturbance (upon PKA phosphorylation) in intact cardiomyocytes.
To further examine the effect of V3599K mutation on the channel gating, independent of CaM, we measured the Ca^{2+} sparks after pretreatment with suramin (10 μM) (followed by washing off), which has been shown to displace endogenous CaM (15–17), in saponin-permeabilized cardiomyocytes. First, without the suramin wash, we examined the effect of exogenously added CaM (100 nM) on the Ca^{2+} spark frequency. With exogenous CaM (100 nM) added in the internal solution, there was no difference in the Ca^{2+} spark frequency in the absence of cAMP among all groups (Figure 8, A and B). However, in the presence of cAMP (1 μM), the Ca^{2+} spark frequency markedly increased in R2474S/+ cardiomyocytes but not in other cardiomyocytes (Figure 8, A and B). In contrast, after treatment with suramin (and without exogenous CaM), namely no CaM condition, the Ca^{2+} sparks were only transiently observed and quickly

Figure 3. Inhibition of ventricular arrhythmia by V3599K mutation in mice. Representative ECG recordings (left) and the summarized data for VT induction (right) after (A) administration of epinephrine (1 mg/kg of body weight, i.p.) or (B) treadmill exercise in WT (+/+)(n = 5), R2474S/+KI (n = 5), V3599K/+KI (n = 5), and R2474S/V3599K-KI mice (n = 5). Representative 24-hour ECG recordings (left) and the summarized data (right) on (C) spontaneous VT or (D) ventricular bigeminy in WT (n = 5), R2474S/+KI (n = 5), V3599K/+KI (n = 5), and R2474S/V3599K-KI mice (n = 5).
disappeared in the absence of cAMP at baseline (data not shown), probably owing to the decrease in intra-SR [Ca^{2+}] caused by the leaky RyR2. However, in the presence of cAMP, the Ca^{2+} spark frequency markedly increased to a similar extent in all groups, without a significant change in SR Ca^{2+} content (Figure 8B). These results clearly indicate that the rescue effect of V3599K mutation on aberrant Ca^{2+} release caused by CPVT-associated mutation (R2474S) is indeed CaM dependent.

CaM binding to the RyR2 was restored in the R2474S/V3599K mice. In resting ventricular cardiomyocytes, only approximately 1% of total CaM is freely diffusible (18), and the bound CaM is highly concentrated along Z lines, where more than 90% of that CaM is RyR2 bound (17). Moreover, we recently found that either angiotensin II or phenylephrine caused translocation of CaM bound to the Z line to the nucleus with G protein–coupled receptor kinase 5, and dantrolene, an RyR2 stabilizer, prevented the translocation of CaM to the nucleus (19), indicating that the reduction in Z line CaM binding is indicative of decreased RyR2 binding. Based on these findings, we examined CaM bound to the Z line along with that in the nucleus in intact cardiomyocytes to assess whether CaM binding to the RyR2 is maintained by V3599K mutation, even in the presence of ISO. In confirmation of our previous report (10), endogenous CaM was well colocalized with the RyR2 on the Z line (Figure 9). The Z line–bound CaM decreased in parallel with
the increase in nuclear CaM only in R2474-KI cardiomyocytes after the addition of ISO, whereas its levels were restored in R2474S/V3599K-KI cardiomyocytes (Figure 9), although the phosphorylation level at Ser2808 was similarly increased by ISO among all mice (Supplemental Figure 6). These data strongly suggest that in response to catecholamine, CaM binding to the RyR2 is restored by the copresence of V3599K mutation upon R2474S-associated channel disorder.

To further examine whether there is any difference in the kinetics of the CaM binding to the RyR2 between WT and KI hearts, we assessed the CaM binding to subcellular fractions by introducing CaM, fluorescently

**Figure 5. Intracellular Ca\(^{2+}\) transients and SR Ca\(^{2+}\) content in intact cardiomyocytes.** (A) Representative recordings of fluo-4 AM fluorescence signal, at a pacing rate of 1 Hz, with ISO (10 nM) activation (ISO) and without it (baseline), and the summarized data (right). n = 28–44 cells from 3 hearts. (B) Representative recording of fluo-4 AM fluorescence signal after addition of caffeine, which is a measure of the SR Ca\(^{2+}\) content, and the summarized data (right). n = 10–19 cells from 3 to 4 hearts. Caffeine-induced Ca\(^{2+}\) transients were measured by first applying a stimulation train at 2 Hz and then by rapidly switching from the superfusing solution to a solution containing 20 mM caffeine for 5 to 6 seconds. Values for individual mice are plotted with mean ± SEM. *P < 0.05, and ***P < 0.001 (1-way ANOVA with post hoc Tukey’s multiple-comparison test).
labeled with Alexa Fluor 488 (Alexa-CaM), into saponin-permeabilized CPVT-KI and WT cardiomyocytes. By this method, we previously confirmed that the fluorescence signal of exogenously introduced Alexa-CaM and that of immunostaining of RyR2 showed a periodical pattern associated with that of sarcomeres, and importantly the staining patterns of both CaM and RyR2 showed an excellent match with each other (Figure 2 in ref. 9). According to the method by Yang et al. (17), we first pretreated the cells with suramin (10 μM) (followed by washing off) to completely abolish endogenous CaM and observed the time course of Alexa-CaM (100 nM) binding along Z lines and determined the maximum specific binding (Bmax) and τ of the curves (Figure 10). Only in R2474S/WT cardiomyocytes, the Bmax of the Alexa-CaM binding decreased in the presence of cAMP, with no significant change in τ. In contrast, there was no difference in the Bmax and τ between WT, R2474S/V3599S, and V3599K/+ cardiomyocytes, regardless of the presence of cAMP. These data consistently support the notion that the CaM-binding affinity to the RyR2 is reduced by R2474S mutation, and V3599K mutation can rescue the reduced CaM-binding affinity.

Direct binding of exogenous CaM to the RyR2 was also evaluated by attaching a UV cross-linker to CaM. Upon cAMP-mediated phosphorylation at Ser2808, the binding affinity of CaM to the RyR2 markedly decreased in R2474S/+KI hearts and was restored in R2474S/V3599K-KI hearts (Figure 11A), although the phosphorylation level at Ser2808 was similarly increased by cAMP among all mice (Supplemental Figure 7). To semiquantitatively evaluate the stoichiometry of CaM binding to RyR2,
we used suramin again to pharmacologically displace CaM from RyR2 (15–17). First, we confirmed the concentration of suramin was enough to completely displace CaM from all 4 subunits of the RyR2. As shown in Figure 11B, suramin (10 μM) almost completely inhibited the exogenous CaM binding to the RyR2 in WT hearts. Then, we observed the inhibitory effect of suramin in KI hearts. Suramin almost completely inhibited the exogenous CaM binding to the RyR2 in all KI mice, regardless of phosphorylation or mutation (Figure 11B and Supplemental Figure 8).

### Discussion

Here, we demonstrated that a single amino acid substitution within the CaM-binding domain of the RyR2, by which the binding affinity of CaM to the RyR2 was markedly enhanced, inhibited Ca\(^{2+}\) leak from the SR, thereby inhibiting CPVT.

Liu et al. recently reported that gene transfer of a mutated CaM (GSH-M37Q), which exhibited a slowed N-terminal Ca dissociation rate and prolonged RyR2 refractoriness, reduced aberrant diastolic Ca\(^{2+}\) release and alleviated arrhythmias in a calsequestrin-associated CPVT model (20). These findings support the important role of CaM in regulating the RyR2 in the pathogenesis of CPVT. However, because CaM interacts with many other proteins in the cell and acts as a regulator or an effector molecule in a variety of cellular functions, it remains to be clarified as to whether direct association of CaM with the RyR2 really modifies the Ca\(^{2+}\) handling in the pathogenesis of CPVT, particularly independent of any possible action of CaM on other proteins. In this regard, the important aspect of this study is that to enhance the binding affinity of CaM, a single amino acid substitution was made only to the CaM-binding site within the RyR2 (V3599K), not to CaM itself. Therefore, without taking any off-target effects into consideration, we can clearly mention that the proper association of CaM to the RyR2 is essential for the prevention of the CPVT phenotype.

One of the major hypotheses to explain the RyR2 dysfunction in heart failure and lethal arrhythmias, such as CPVT, is that defective interaction between the N-terminal (N: 1–600) and the central (C: 2000–2500) domains (4, 5) causes an instability of the channel. Namely, in the resting state, the N-terminal and central RyR2 domains interact with each other to act as a regulatory switch that influences RyR channel gating. This tight interdomain interaction, termed “domain zipping,” seems to stabilize the closed channel. Weakening of these interdomain interactions may be caused by mutations in either the N-terminal or central regions of RyR2. In support of this idea, using 3-dimensional cryoelectron microscopy, it has been reported that the N-terminal regions within domain 5 are closely located with the central regions within domain 6 (21), that the interaction between the N-terminal and central mutation regions is an intersubunit (not intrasubunit) interaction (22), and that the 3-dimensional location of the Ser2808 PKA phosphorylation site was located near the domain-domain boundary (23), suggesting that PKA hyperphosphorylation at Ser2808 may alter the domain-domain interactions between the N-terminal and the central regions of the RyR2 and thus the gating properties of the channel. As a matter of fact, we previously reported that PKA phosphorylation at Ser2808 induced domain unzipping and aberrant Ca\(^{2+}\) release only in CPVT-associated R2474S/+KI (but not WT) mice (6). Furthermore, accumulated evidence has recently indicated that domain unzipping is allosterically linked to reduced affinity of CaM

---

**Table 2. Effect of isoproterenol on Ca\(^{2+}\) spark characteristics in intact cardiomyocytes**

<table>
<thead>
<tr>
<th>ISO (10 nM)</th>
<th>Peak (F/F(_{0}))</th>
<th>FWHM (μm)</th>
<th>FDHM (ms)</th>
<th>No. of sparks</th>
</tr>
</thead>
<tbody>
<tr>
<td>WT (+/+) (n = 3)</td>
<td>1.20 ± 0.02</td>
<td>1.53 ± 0.15</td>
<td>36.44 ± 4.85</td>
<td>38</td>
</tr>
<tr>
<td>R2474S/+ (n = 3)</td>
<td>1.19 ± 0.01</td>
<td>1.53 ± 0.10</td>
<td>36.17 ± 4.36</td>
<td>32</td>
</tr>
<tr>
<td>V3599K/+ (n = 3)</td>
<td>1.17 ± 0.01</td>
<td>1.43 ± 0.11</td>
<td>40.21 ± 6.65</td>
<td>31</td>
</tr>
<tr>
<td>WT (+/+) (n = 3)</td>
<td>1.18 ± 0.01</td>
<td>1.67 ± 0.15</td>
<td>34.02 ± 3.22</td>
<td>35</td>
</tr>
<tr>
<td>R2474S/+ (n = 3)</td>
<td>1.25 ± 0.01</td>
<td>1.81 ± 0.09</td>
<td>36.46 ± 2.89</td>
<td>65</td>
</tr>
<tr>
<td>R2474S/V3599K (n = 3)</td>
<td>1.26 ± 0.01</td>
<td>2.33 ± 0.06*</td>
<td>81.14 ± 3.49*</td>
<td>352</td>
</tr>
<tr>
<td>V3599K/+ (n = 3)</td>
<td>1.24 ± 0.02</td>
<td>2.21 ± 0.06*</td>
<td>53.93 ± 2.84*</td>
<td>211</td>
</tr>
</tbody>
</table>

*P < 0.01 vs. WT (+/+) with ISO. \(P < 0.05\) vs. WT (+/+) with ISO. \(^\dagger\)P < 0.01 vs. R2474S/+ with ISO. \(^\ddagger\)P < 0.05 vs. R2474S/+ with ISO. \(n\), the number of hearts; FWHM, full width at half maximum; FDHM, full duration at half maximum.
bound to the RyR2, thereby causing aberrant diastolic Ca\textsuperscript{2+} release as a common pathogenic mechanism underlying CPVT and heart failure (10–13, 24–26).

On the basis of the above-mentioned structural background regarding the pathogenic mechanism of RyR2-associated CPVT, we propose a hypothetical model to explain why CaM was not displaced from the R2474S-mutated subunits by a single amino acid substitution (V3599K) applied against the subunits not harboring R2474S mutation (Figure 12). Because the difference in the extent of CaM binding between WT or R2474S/V3599K-KI hearts and the R2474S/+-KI hearts (with cAMP) was approximately half of that in the presence and absence of suramin in WT hearts (Figure 11B), it is suggested that in R2474S/+-KI hearts the defective intersubunit interaction, namely domain unzipping, that occurred in the R2474S-mutated subunits may allosterically decrease the binding affinity of CaM to the R2474S-mutated subunits and in turn displace approximately half of the bound CaM from the RyR2. On the other hand, in R2474S/V3599-KI hearts the defective intersubunit interaction would be minimized owing to the strong binding of CaM to the V3599K-mutated subunits, thereby preventing the decrease in the binding affinity of CaM to the remaining R2474S-mutated subunits and subsequent displacement of CaM from the RyR2. This postulated model is supported by the previous finding obtained by fluorescence resonance energy transfer method that the domain switch and CaM-binding regions are distinct but interact allosterically (rather than orthosterically) (26).

Dantrolene, which directly binds to the N-terminal domain (amino acid residues 600–610) of the RyR2, restored the domain zipping state in catecholamine-added R2474S/+-KI and failing hearts and induced the reassociation of CaM to the RyR2, thereby inhibiting Ca\textsuperscript{2+} leak, DaD, and lethal arrhythmia (6, 9, 11, 27). We previously showed using a CPVT-type–KI mouse model that a single amino acid mutation (N4104K) within the CaM-like domain (CaMLD) observed in patients with CPVT causes an abnormally tight interaction between the CaM-like domain (CaMLD) and the CaM-binding domains (CaMBDs), thereby decreasing the affinity of CaM to CaMBDs and in turn causing Ca\textsuperscript{2+} leak and lethal arrhythmia (28). In this KI mouse model, dantrolene did not inhibit the Ca\textsuperscript{2+} leak, but the increase in the affinity of CaM to

Figure 7. SCaTs in intact cardiomyocytes. (Left) Representative traces of SCaTs and (right) the summarized data as percentage occurrence of SCaTs in 874 to 1693 cells from 3 to 4 hearts in isolated intact cardiomyocytes. Arrows indicate SCaTs. The SCaTs were induced after cessation of pacing at 1, 2, 3, 4, and 5 Hz of stimulation with or without 10 nM ISO. SCaT incidence was much higher in R2474S/+-KI cells than in WT or V3599K/+-KI cells in the presence of the ISO (10 nM). However, SCaT incidence was markedly decreased in R2474S/V3599K-KI cells. Values for individual mice are plotted with mean ± SEM. *P < 0.05, **P < 0.01, and ***P < 0.001 (1-way ANOVA with post hoc Tukey’s multiple-comparison test).
the RyR (GSH-CaM) inhibited it (28). This notion supports the ideas that dissociation of CaM from the RyR2, induced either by defective N-terminal and central domain interaction (unzipping) or by defective CaMLD and CaMBD interaction, is a common abnormal pathway leading to Ca\(^{2+}\) leak and that the therapeutic effect of dantrolene on the aberrant Ca\(^{2+}\) release is indeed CaM dependent. In support of this idea, Oo et al. recently showed that in single-channel recording, dantrolene caused inhibition of the RyR1 and RyR2 only after adding physiological CaM (100 nM) but not in the absence of CaM (29).

There is a limitation of the study. It was not feasible to create double mutations (R2474S and V3599K) in the same subunit of the RyR2 by natural mating. Therefore, further study is needed to clarify whether the double mutations in the same subunit indeed restore the stability of the channel.

**Figure 8. Ca\(^{2+}\) sparks in saponin-permeabilized cardiomyocytes.** (A) Representative recordings of line-scan images of fluo-4 AM fluorescence (left) in the presence of exogenous CaM (100 nM) with or without cAMP (1 μM) or (right) with cAMP (1 μM) plus suramin (10 μM). (B) Summarized data for Ca\(^{2+}\) spark frequency and SR Ca\(^{2+}\) content. To measure SR Ca\(^{2+}\) content, caffeine-induced Ca\(^{2+}\) transients were recorded by adding 20 mM caffeine. Values for individual mice are plotted with mean ± SEM. *P < 0.05, **P < 0.01, and ***P < 0.001 (1-way ANOVA with post hoc Tukey’s multiple-comparison test); n = 27–42 cells from 3 to 5 hearts.
In conclusion, enhancement of the CaM-binding affinity of the RyR2 is essential to prevent CPVT-associated arrhythmogenesis. Thus, CaM may act as an intrinsic natural stabilizer for the RyR2 channel function, and hence increasing the affinity of CaM to the RyR2 could be a potent therapeutic target for heart failure and lethal arrhythmia.

Figure 9. Localization and the binding characteristics of the endogenous CaM in intact WT and KI cardiomyocytes. (A) Representative images of the endogenous CaM, which is colocalized with the RyR2 in WT cardiomyocytes, in the (left) absence or (right) presence of ISO (10 nM); immunostaining of CaM (shown in green); immunostaining of the RyR2 (red). Scale bars: 10 μm. (B) Summarized data of the Z line–bound CaM and the nuclear CaM. The immunofluorescence signal of the Z line–bound CaM was divided by that of the RyR2, normalized to control (baseline of WT), and expressed as a ratio. The immunofluorescence signal of the nuclear CaM was divided by that of DAPI for nuclear staining, normalized to control (baseline of WT), and expressed as a ratio. Data are presented as mean ± SEM of 37 to 50 cells from 4 hearts. *P < 0.05 (1-way ANOVA with post hoc Tukey’s multiple-comparison test).
Methods

Animals. C57BL/6 mice, 20–25 weeks old, were used in this study. WT C57BL/6 mice were obtained from Japan SLC Inc. V3599K-KI mice were from UNITECH Co. Ltd. Mice were assigned to each experiment randomly, and the order of the group in each experiment was decided randomly. Allocation concealment was not applicable because the genetic background of each experimental group was different and this study did not involve any testing of therapeutic agents. Measurement and analysis of the obtained data were performed by the investigators, who were blinded to the genetic information of the mice. This study conformed to the Guide for the Care and Use of Laboratory Animals published by the National Academies Press (revised 1996). The care of the animals and the protocols used were in accordance with guidelines laid down by the Animal Ethics Committee of Yamaguchi University Graduate School of Medicine.

Echocardiogram. Cardiac function was analyzed by an F37 ultrasound machine (Hitachi medical) equipped with a 7.5-MHz probe (Hitachi, UST-5413). Mice were initially anesthetized with 4% to 5% isoflurane (mixed with oxygen) and maintained with 1% to 2% isoflurane during echocardiography.
Surface ECG. ECG was monitored for the WT and the series of KI mice using ECG telemetry. Briefly, transmitters (Data Sciences International) were implanted in the back space with subcutaneous electrodes in a lead II configuration. Telemetry was first recorded with mice in a conscious state for 24 hours. After that, ECG was recorded at baseline and after the injection of epinephrine (1 mg/kg of body weight, i.p.) and monitored for 30 minutes in a subset of WT, R2474S/+-KI, V3599K/+-KI, and R2474S/V3599K-KI mice. Another set of WT, R2474S/+-KI, V3599K/+-KI, and R2474S/V3599K-KI mice were subjected to a treadmill exercise activity (Panlab).

Figure 11. CaM binding to the RyR2 in cardiac homogenates. (A) (Top) Representative immunoblots of the RyR2-bound CaM-SANPAH (a photoreactive cross-linker). The WT and KI cardiac homogenates were first diluted in binding buffer (150 mM KCl, 10 μM CaCl, and 20 mM MES at pH 6.8) and then reacted with various concentrations of CaM-SANPAH (32–1024 nM) in the presence or absence of cAMP (1 μM) and okadaic acid (1 μM). CaM binding to the RyR2 was determined by immunoblotting with anti-CaM antibody to detect the RyR2-bound CaM. (Bottom) Summarized data of CaM binding to the RyR2 as a function of the concentration of CaM-SANPAH. CaM binding was expressed as the ratio to the maximum binding of CaM (at 1024 nM). Data are presented mean ± SEM of 3 hearts. (B) Effect of cAMP (1 μM) and suramin (10 μM) on CaM binding to the RyR2. The concentration of CaM-SANPAH was fixed at 128 nM. Data are presented as mean ± SEM of 3 hearts. *P < 0.05; NS, not significant (Student’s t test).
Isolation of cardiomyocytes. Cardiomyocytes were isolated from the mouse hearts as described previously (6). Briefly, mice were anesthetized with pentobarbital sodium (70 mg/kg of body weight, i.p.), intubated, and ventilated with ambient air. An incision in the chest was made, and the heart was quickly removed and retrogradely perfused with a collagenase-free buffer (Thermo Fisher Scientific, DMEM, 10313) via the aorta under constant flow. The left ventricular myocardium was minced with scissors in a fresh collagenase-containing buffer, and the rod-shaped adult mouse cardiomyocytes were prepared by retrograde perfusion of the hearts with 95% O2 and 5% CO2-bubbled minimal essential medium (MilliporeSigma) supplemented with 50 μM [Ca2+], 0.5 mg/ml collagenase B (Roche), 0.5 mg/ml collagenase D (Roche), and 0.02 mg/ml protease type XIV (MilliporeSigma, P-5147). The Ca2+ concentration was then gradually increased to a final concentration of 1 mM by changing the incubation medium (50, 100, 300, and 600 μM and then 1 mM). The isolated mouse cardiomyocytes were transferred to laminin-coated glass culture dishes and incubated for a few hours at 37°C in a 5% CO2 and 95% O2 atmosphere.

Monitoring of Ca2+ transients of cardiomyocytes. Isolated ventricular myocytes were incubated with 20 μM fluo-4 AM (Molecular Probes) for 20 minutes at room temperature and washed twice with Tyrode’s solution (made in-house; 140 mM NaCl, 1 mM MgCl2, 10 mM glucose, 6 mM KCl, 5 mM HEPES pH7.4, 2 mM CaCl2) as described previously (6). All experiments were conducted at 35°C. Intracellular Ca2+ measurements with cells stimulated by a field electric stimulator (Ion Optix) were taken using a fluorescent digital microscope (Keyence, BZ9000). The relative occurrence of spontaneous Ca2+ release upon cessation of stimulation at 1, 2, 3, 4, and 5 Hz was measured in WT and KI myocytes.

Analysis of Ca2+ sparks and SR Ca2+ content. The Ca2+ sparks were measured as previously described (6) using a laser-scanning confocal microscope (LSM-510, Carl Zeiss) equipped with an argon ion laser coupled with an inverted microscope (Axiovert 100, Carl Zeiss) with a Zeiss ×40 oil immersion Plan-Neofluor objective (numerical aperture, 1.3; excitation at 488 nm; emission at >505 nm). Briefly, intact cardiomyocytes were loaded with fluo-4 AM (20 μM) for 20 minutes at 24°C. Line-scan mode was used, where a single cardiomyocyte was scanned repeatedly along a line parallel to the longitudinal axis, avoiding nuclei. To monitor Ca2+ sparks, cardiomyocytes were stimulated until the Ca2+ transients reached a steady state. Stimulation was then stopped, and Ca2+ sparks were recorded during the subsequent approximately 10 seconds’ rest. Data were analyzed with SparkMaster (https://sites.google.com/site/sparkmasterhome/), an automated analysis program that allows for rapid and reliable spark analysis. The variables analyzed included general image parameters (like number of detected sparks, spark frequency) as well as individual spark parameters (amplitude, FWHM, and FDHM). To assess the SR Ca2+ content, caffeine (10 mM) was rapidly perfused to discharge the SR-loaded Ca2+.
Immunocytochemistry analysis of endogenous RyR2-bound CaM. Isolated cardiomyocytes were fixed with 4% paraformaldehyde (WAKO, 163-20145) in PBS for 5 minutes, washed 3 times with PBS, and permeabilized in 0.5% Triton X-100 (MilliporeSigma) and 1% BSA (Nacalai Tesque) for 20 minutes. Then, the cardiomyocytes were incubated overnight at 4°C with the anti-CaM antibody (EP799Y, Abcam) and anti-RyR antibody (MilliporeSigma, C3-33) in 1% BSA and 0.5% Triton X-100, followed by labeling with an Alexa Fluor 488–conjugated goat anti-rabbit (Thermo Fisher Scientific, A11008) and an Alexa Fluor 633–conjugated goat anti-mouse (Thermo Fisher Scientific, A21052) secondary antibody. The cardiomyocytes were washed 3 times with PBS.

The sarcomere-related periodical increase in the Alexa Fluor 633 and Alexa Fluor 488 fluorescence intensity from baseline was integrated with the longitudinally selected distance (~25 μm), and then the value was divided by the distance. Then mean value of 1 sarcomere-related increased fluorescence intensity was calculated as the arbitrary amount of RyR and RyR-bound CaM. Secondary antibody labeling alone showed no detectable fluorescence pattern.

Determination of the binding of exogenous CaM to the RyR2 in saponin-permeabilized cardiomyocytes. The exogenous CaM, fluorescently labeled with Alexa Fluor 633 (Molecular Probes), was introduced into the saponin-permeabilized WT and KI cardiomyocytes as described previously (10). Then, the distribution of localized CaM was determined by densitometric measurement of Alexa-CaM fluorescence. Briefly, the fluorescently labeled cardiomyocytes were laser-scanned with the confocal microscope (LSM-510, Carl Zeiss) (numerical aperture, 1.3; excitation at 633 nm; emission 650 nm). The sarcomere-related periodical increase in the Alexa Fluor 633 fluorescence intensity from baseline was integrated with the longitudinally selected distance (~25 μm), and then we divided the value by the distance.

Analysis of the binding characteristics of CaMBPs to CaM and the association of CaM to the RyR2 by CaM-SANPAH cross-linking method. The binding of CaM to the RyR2 was evaluated using the photoreactive cross-linker sulfo-SANPAH (Thermo Fisher Scientific) as described previously (10). First, we made a CaM-SANPAH conjugate by mixing 50 μM recombinant CaM in the conjugation buffer (150 mM KCl and 20 mM MOPS at pH 7.2) and 100 μM sulfo-SANPAH in the dark. After 30 minutes, conjugation was quenched by adding excess amount of lysine. CaM-SANPAH conjugate was purified using Amicon Ultra (MW cutoff 10 kDa). Canine SR vesicles (for analysis of the binding characteristics of CaMBPs to CaM) and mouse cardiac homogenates (for analysis of association of CaM to the RyR2) were diluted with the binding buffer (150 mM KCl, 10 μM CaCl₂, and 20 mM MES at pH 6.8) to 1 mg/ml and mixed with 100 nM CaM-SANPAH conjugate in the dark in a glass tube with or without CaMBPs. After 10 minutes’ binding time, UV irradiation was done for cross-linking. Then sample buffer was added to the cross-linked SR membrane, and Western blotting using anti-CaM antibody (Merck, MilliporeSigma) was done. CaM-SANPAH cross-linked to the RyR2 was detected with a 550-kDa band.

The binding affinity of CaM to each CaMBP was measured as an inhibition of CaM-SANPAH binding to the RyR2 in SR vesicles. When the affinity of CaM to CaMBP is higher than to the RyR2, most of the added CaM-SANPAH binds to CaMBP instead of the RyR2. To assess the conformational state of CaM, we measured the fluorescence level of tryptophan within CaM upon its binding to CaMBPs.

CaMBP-CaM interaction measurement by QCM. Binding of CaM to WT CaMBP or V3599K CaMBP was detected by using a 27-MHz QCM (Initium Inc.), a highly sensitive mass-measuring apparatus, as we described previously (8). The QCM Au electrode was coated with CaMBP and immersed in a solution (500 μl) containing 150 mM NaCl, 20 mM MOPS, and 0.01 mM Ca²⁺ (pH 7.2). The amount of peptide binding was determined from the frequency changes due to changes in mass on the electrode (with sensitivity of the order of subnanogram) on injection of a small volume (2–5 μl) of solution containing CaM. No significant nonspecific binding of CaM to the Au electrode was detected.

Histology. Hearts from the WT and the series of KI mice, aged 16–22 weeks, were fixed using 10% formalin. A complete, full-circumferential section, at the level of the 2 left ventricular papillary muscles, was selected for morphometric analysis. Hematoxylin and eosin and Masson’s trichrome stains were performed for each section of the ventricle.

Immunoblot analysis. We performed immunoblot analysis of RyR2 (C3-33, MilliporeSigma), phosphorylated RyR2 at Ser2808 (p–Ser2808 RyR2), SR Ca²⁺-ATPase (SERCA2a, Affinity Bioreagents), phospho-amban (PLB, Santa Cruz Biotechnology Inc.), phosphorylated PLB at serine 16 (p–Ser16 PLB, Fitzgerald), phosphorylated PLB at threonine 17 (p–Thr17 PLB, Santa Cruz Biotechnology Inc.), and glyceraldehyde-3-phosphate dehydrogenase (MilliporeSigma) as described previously (3).
Statistics. Two-tailed Student’s *t* tests were used for statistical comparisons of data obtained during the 2 different situations, whereas 1-way ANOVA with a post hoc Tukey’s test was used for statistical comparison of more than 2 groups. All data are expressed as mean ± SEM. A *P* value of less than 0.05 was considered statistically significant.

Study approval. All experiments were approved by the Animal Ethics Committee of Yamaguchi University Graduate School of Medicine.

Author contributions
YN, TY, and SK designed the research, performed experiments, analyzed data, and wrote the manuscript. YH and MT conducted the in vivo ECG recording and provocation test for VT; GF and TK assisted with the measurement of Ca²⁺ transients, cell shortening, and Ca²⁺ sparks and data analysis thereof. XX and SN assisted with Western blot experiments. HU, TO, and SO assisted with CaM-binding assays and data analysis thereof. MY designed and supervised the research and wrote the manuscript.

Acknowledgments
This work was supported by grants-in-aid for scientific research from the Ministry of Education, Culture, Sports, Science and Technology in Japan (17H04178 and 26293189 to MY, 15K09085 to TY, 15K09142 and 18K08039 to SK). We wish to acknowledge Yoko Okamoto and Satomi Tateda for their technical assistance.

Address for correspondence: Masafumi Yano, Department of Medicine and Clinical Science, Division of Cardiology, Yamaguchi University Graduate School of Medicine, 1-1-1 Minamikogushi, Ube, Yamaguchi 755-8505, Japan. Phone: 81.836.22.2248; Email: yanoma@yamaguchi-u.ac.jp.


insight.jci.org   https://doi.org/10.1172/jci.insight.126112
21. Wang R, et al. Localization of an NH(2)-terminal disease-causing mutation hot spot to the “clamp” region in the three-dimen-
22. Liu Z, et al. Dynamic, inter-subunit interactions between the N-terminal and central mutation regions of cardiac ryanodine
25. Oda T, et al. Oxidation of ryanodine receptor (RyR) and calmodulin enhance Ca release and pathologically alter, RyR structure
26. Oda T, et al. In cardiomyocytes, binding of unzipping peptide activates ryanodine receptor 2 and reciprocally inhibits calmod-
27. Kobayashi S, et al. Dantrone, a therapeutic agent for malignant hyperthermia, inhibits catecholaminergic polymorphic ventric-
28. Nishimura S, et al. Mutation-linked, excessively tight interaction between the calmodulin binding domain and the C-terminal
domain of the cardiac ryanodine receptor as a novel cause of catecholaminergic polymorphic ventricular tachycardia. *Heart