Cypher/ZASP Is a Novel A-kinase Anchoring Protein*

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Background: Cypher/ZASP plays an essential structural role in cardiac muscle.
Results: Cypher/ZASP specifically interacted with PKARIIα and calcineurin.
Conclusion: Cypher/ZASP is a novel AKAP acting as a sarcomeric signaling center for potential phosphorylation regulation of the function of channels and myofilament proteins.
Significance: Cypher/ZASP-PKA-calcineurin complex expands our understanding the role of Cypher/ZASP in the heart.

PKA signaling is important for the post-translational modification of proteins, especially those in cardiomyocytes involved in cardiac excitation-contraction coupling. PKA activity is spatially and temporally regulated through compartmentalization by protein kinase A anchoring proteins. Cypher/ZASP, a member of PDZ-LIM domain protein family, is a cytoskeletal protein that forms multiprotein complexes at sarcomeric Z-lines. It has been demonstrated that Cypher/ZASP plays a pivotal structural role in the structural integrity of sarcomeres, and several of its mutations are associated with myopathies including dilated cardiomyopathy. Here we show that Cypher/ZASP interacts with the L-type calcium channel through its C-terminal PDZ binding motif. Expression of Cypher/ZASP facilitated PKA-mediated phosphorylation of the L-type calcium channel in vitro. Additionally, the phosphorylation of the L-type calcium channel at Ser1928 induced by isoproterenol was impaired in neonatal Cypher/ZASP-null cardiomyocytes. Moreover, Cypher/ZASP interacted with the Ser/Thr phosphatase calcineurin, which is a phosphatase for the L-type calcium channel. Taken together, our data strongly suggest that Cypher/ZASP not only plays a structural role for the sarcomeric integrity, but is also an important sarcomeric signaling scaffold in regulating the phosphorylation of channels or contractile proteins.

In cardiomyocytes, the cAMP/PKA signaling initiated by β-adrenergic receptors (β-ARs)3 is of central importance for all functions, including excitation-contraction coupling, metabolism, hypertrophy, and survival (1, 2). The PKA holoenzyme is a heterotetramer containing two catalytic (C) and two regulatory (R) subunits (3). Four genes encode PKA regulatory subunits (RIα, RIβ, RIα, and RIβ) in mammals. Binding of the second-messenger cAMP to the regulatory subunits releases the catalytic subunits of PKA, which catalyze its substrate to be phosphorylated at either the serine or the threonine within the consensus sequence motif (RXS/T)X, X is variable) (4). The regulatory subunits also regulate the cellular localization of PKA by binding to a special group of proteins: A-kinase anchoring proteins (AKAPs).

AKAPs spatially and temporally restrict or compartmentalize the activity of PKA. To date, ~70 AKAP genes have been identified, among which ~20 are expressed in the heart (5). Three types of AKAPs have been classified. Type II AKAPs specifically bind to PKA RIα, whereas type I AKAPs bind to RI. A few AKAPs with dual specificity bind to both RI and RI3.

PKA-mediated phosphorylation of sarcomeric proteins induced by β-AR stimulation, including cardiac troponin I (cTnI) (6), myosin-binding protein C (MyBP-C) (7), titin (8), and myosin light chain (9), is also important for cardiac contraction and left ventricular torsion. The phosphorylation of cTnI and cardiac MyBP-C leads to decreased calcium responsiveness, thus increasing the myofibril relaxation rate. Some cardiac AKAPs have been shown to localize at sarcomeres, such

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as synemin (10), cardiac troponin T (11), myospryn (12), and myomegalin (13). Synemin and myospryn co-localize with PKARII at the Z-line or the Z-line/costamere in striated muscle. Myospryn also interacts with calcineurin (CaN) (14). Myomegalin might be an AKAP for the sacromeric proteins MyBP-C and cTnI. Cardiac troponin T is a dual specificity AKAP regulating cTnI phosphorylation through the tropoerin complex.

Initially, AKAPs were regarded as recruiters of PKA and phosphatases to form a signaling complex for each of its unique substrates. Recently, AKAP complexes have also been reported to regulate gene transcriptional expression. A direct role of AKAP79/150 has been suggested through its organized signal complexes cAMP/CREB (cAMP-response element-binding protein) or CaN/NFAT (15, 16).

Cypher/ZASP is a striated Z-line protein, which plays an important structural role in cardiac muscle in maintaining the integrity of sarcomeres under the stress of contraction force (17–20). Here, we report that the Z-line protein Cypher/ZASP is also a typical type II AKAP that specifically interacts with the RI\(\alpha\) regulatory subunit of PKA and the Ser/Thr phosphatase CaN, making Cypher/ZASP-PKA-CaN a signaling center for sarcomeric proteins or channels such as the L-type calcium channel (LTCC).

EXPERIMENTAL PROCEDURES

Antibodies and Mice—FLAG epitope, \(\alpha\)-actinin, and plakoglobin antibodies were from Sigma-Aldrich. GST and Myc epitope antibodies were from Abcam. Antibodies against PKA-c, p-Erk1/2, Erk1/2, CaN, and PKA substrate were from Cell Signaling. LTCC phospho-Ser\(^{1928}\) antibody was from Badrilla Ltd. LTCC and GAPDH antibodies were from Santa Cruz Biotechnology. PKA RI\(\alpha\) antibody was either from Millipore or from Abcam. Calmodulin antibody was from Assay Biotech. The rabbit polyclonal Cypher antibody was generated by us.

Generation of Cypher knock-out mice has been described previously (19). Mice were maintained in a pathogen-free vivarium, and all procedures involving mice were approved by the Institutional Animal Care and Use Committee of Zhejiang University.

Plasmids—Plasmids containing the coding sequences for RI\(\alpha\), RI\(\beta\), RI\(\alpha\), and RI\(\beta\) were gifts from Dr. Susan S. Taylor (University of California San Diego, La Jolla, CA). All tagged expression vectors (GST-, FLAG-, Myc-) were constructed in the pXJ40 vector as described previously (21). KOD polymerase was used for amplification, and DNA sequences were confirmed by DNA sequencing.

Protein-Protein Interactions in Vitro—Protein-protein interactions were studied in vitro using overexpression of tagged proteins in HEK293 cells. Plasmids were co-transfected using Lipofectamine 2000 (Invitrogen). 36 h later, cells were harvested and resuspended in radioimmune precipitation buffer assay (50 mm Tris, pH 7.4, 150 mm NaCl, 1% Nonidet P-40, 0.5% sodium deoxycholate, 0.1% SDS) with protease inhibitors (Roche Applied Science). To analyze LTCC phosphorylation, transient transfected cells were incubated with 100 \(\mu\)M forskolin (Sigma), 250 \(\mu\)M isobutylmethylxanthine for 30 min before harvesting the cells. Co-immunoprecipitation assays were performed using anti-protein tag antibodies and protein A-agarose beads (Roche Applied Science). Binding proteins were further analyzed by SDS-PAGE and immunoblot.

Immunostaining—3-month-old male mice were injected with isoproterenol (15 mg/kg, intraperitoneal). Saline was injected as control. 30 min later, mice were anesthetized, and hearts were dissected, immersed in iso-pentane, and snap-frozen for cryosectioning (Leica CM3050 S).

Immunostaining of frozen sections (12 \(\mu\)m) was done as described (22). The samples were fixed in ice-cold acetone for 5 min, rehydrated in PBS, and incubated with primary antibody in gold buffer (155 mm NaCl, 2 mm EGTA, 2 mm MgCl\(_2\), 20 mm Tris-HCl, pH 7.5) overnight at 4 °C. After washing out primary antibody by PBS, the samples were incubated with secondary antibodies and DAPI at room temperature for 1 h, washed with PBS, and embedded in fluorescent embedding medium (DAKO). Images were taken using a Leica SP5 confocal microscope equipped with a 63 \(\times\) glycerol immersion objective in sequential scanning mode.

Neonatal Rat Cardiomyocyte Isolation—Neonatal rat cardiomyocytes were isolated from 1-day-old rats as described previously (23). Lentivirus-mediated gene transfer was performed 12 h later followed by 24 h in culture before harvesting the cells for protein analysis.

Structure Modeling—The PKARII-Cypher AA202–215 structure was modeled using Xtalview (24) based on the crystal structure of the RII-D/D-AKAP-JS complex (Protein Data Bank (PDB) accession code: 2izx) (25). The modeled structure was further refined using REFMAC (26). All structural illustrations were prepared with PYMOL (52). The model showed that Thr\(^{203}\) from the Cypher peptide formed van der Waals contacts with Ile\(^{3}\), Ile\(^{5}\), and Ala\(^{24}\) from RII dimerization/docking domain (D/D). The substitution of Thr\(^{203}\) to Ile\(^{203}\) enhanced the van der Waals interactions.

CaN Enzymatic Activity Measurement—The CaN activity assay was performed as described (27). The CaN enzymatic activity was measured by using a colorimetric assay kit according to the manufacturer’s instructions (Nanjing Jiancheng Bioengineering Institute, Nanjing, China) (28). The free phosphate released from synthetic RII phosphopeptide substrate by calcineurin was measured using malachite green assay. Cyclosporin A, a specific inhibitor for calcineurin, was used. The calcineurin specific activity was calculated from subtracting the value measured in the presence of cyclosporin A from total activity measured without cyclosporin A.

To measure activated calcineurin, calcineurin/calmodulin co-immunoprecipitations were performed as described (27, 29). Mouse hearts were lysed in co-immunoprecipitation buffer (20 mmol/liter NaPO\(_4\), 150 mmol/liter NaCl, 2 mmol/liter MgCl\(_2\), 0.1% Nonidet P-40, 10% glycerol, 1 mmol/liter DTT) with protease inhibitor (Roche Applied Science) and phosphatase inhibitor (Roche Applied Science). The lysate was incubated with calcineurin or calmodulin antibody followed by the addition of protein A-agarose beads. The washed beads were subjected to SDS-PAGE and immunoblot.

Real-time PCR—Total RNA was isolated from 1-day-old mouse hearts using TRIzol reagent (Invitrogen) according to
the manufacturer’s protocol. Real-time PCR was performed as described previously (30). The oligonucleotide primers for MCIP1.4 were: forward: 5’-TCAGCTGGGTTACGGAG-3’, and reverse, 5’-ACTGGAAGTTGGTTGCCTTGTC-3’.

Statistics—Data were expressed as mean ± S.E. We performed statistical evaluation using Student’s unpaired t test. \( p < 0.05 \) was considered to be statistically significant.

RESULTS

Cypher/ZASP Interacted with the Regulatory Subunit RII\(\alpha\) of PKA—Cypher/ZASP contains a PDZ domain at its N terminus and three LIM domains at its C terminus with a short ZASP motif (ZM, AA186–211) in the middle. By analyzing the secondary structure of the Cypher1c amino acid sequence (AF114378_1), we found that the fragment AA200–217 could potentially form an amphipathic helix (Fig. 1A), a feature of AKAPs. The AA200–217 sequence partly overlaps with the ZM region (AA186–211). To determine whether Cypher indeed interacts with PKA regulatory subunits through its ZM domain, GST-tagged Cypher was co-expressed with each of the four AKAPs. The AA200–217 sequence partly overlaps with the ZM region (CCSR, AA108–227), being encoded by the exon 4, interacted with the PKA RII\(\alpha\) (Fig. 1, A, C, and D). To further determine whether the \(\alpha\) helix was essential for the interaction, we substituted proline at each of the following amino acid residue positions, Thr203, Leu204, and Leu215, to disrupt the helix. All three variants (T203P, L204P, and L215P) significantly decreased the binding of Cypher to PKARII\(\alpha\) (Fig. 1E), whereas the mutant Cypher with an AA200–217 deletion had no interaction (Fig. 1F). Therefore, the CCSR of Cypher specifically interacted with PKA RII\(\alpha\).

The D/D of RII\(\alpha\) is involved in the interactions of typical AKAPs. To further study whether the D/D domain was involved in the interaction with Cypher, FLAG-tagged Cypher1c was co-expressed with either the GST-tagged full-length, D/D domain (1–45) or the D/D deletion (\(\Delta D/D\)) of RII\(\alpha\) in HEK293 cells. Both full-length and D/D domain interacted with Cypher, whereas \(\Delta D/D\) did not (Fig. 1G). Therefore, Cypher interacted with the D/D region of RII\(\alpha\), displaying the characteristics of a genuine AKAP.

Cypher belongs to the ALP/Enigma subfamily of the PDZ-LIM domain protein family. We further investigated the interactions between PKA RII\(\alpha\) and other members of the ALP/Enigma subfamily. We found that, like Cypher, enigma homolog (ENH) bound to PKA RII\(\alpha\), whereas ALP and CLP36 did not (Fig. 1H).

To confirm the interaction in vivo, an anti-Cypher antibody was used for a co-immunoprecipitation assay. RII protein was pulled down together with Cypher (Fig. 1I) in wild-type but not Cypher-null mouse hearts. Rabbit IgG failed to pull down either Cypher or PKARII.

To further verify the interaction between Cypher and PKARII in cardiomyocytes, immunostaining was performed (Fig. 2). As shown previously (17), Cypher localized predominantly at Z-lines together with sarcomeric \(\alpha\)-actinin (Fig. 2A). Isoproterenol treatment did not disturb the Z-line localization of Cypher (Fig. 2B); however, it lead to more pronounced localization of Cypher at intercalated disks (Fig. 2B, inset). PKARII localized at intercalated disks but did not show significant striated staining pattern in control hearts (Fig. 2C). However, hearts treated with isoproterenol showed increasing localization of PKARII at Z-lines, as judged by colocalization with \(\alpha\)-actinin (Fig. 2D). Thus, endogenously expressed Cypher and PKARII may also form a protein complex at Z-lines of mouse cardiomyocytes.

The Cardiomyopathy-associated Mutation T206I (T203I in Mouse) Enhanced the Interaction between ZASP and PKARII—Several ZASP mutations have been identified in human patients with myopathies including dilated cardiomyopathy. Some of the mutations are localized to the sequence encoded by exon 4. Similar to Cypher CCSR, the peptide encoded by the ZASP exon 4 interacted with PKA RII\(\alpha\) (Fig. 3A). Mutation S189I did not change the interaction, whereas mutation T206I significantly enhanced it (Fig. 3A). For further investigation of the interaction, the RII\(\alpha\)-AKAP-IS crystal structure was adapted, and the AKAP-IS sequence was superimposed over the Cypher sequence (AA202–215). As shown in the model, the Thr\(^{203}\) residue forms van der Waals contacts with the Ala\(^{21}\), Ile\(^{3}\), and Ile\(^{5}\) residues of RII D/D, whereas substitution of Thr\(^{203}\) to Ile\(^{203}\) enhances the van der Waals interactions (Fig. 3B). Surface representations of the Cypher helix (AA202–215) showed that the substitution of Thr\(^{203}\) to Ile\(^{203}\) also increased the hydrophobic surface involving the PKA RII D/D interaction (Fig. 3C).

Cypher/ZASP Facilitated Phosphorylation of the LTCC by PKA—It has been reported that the LTCC contains a PDZ-binding motif at its cytosolic tail and interacts with the PDZ domain of ENH (31), which together with Cypher and Enigma, belongs to the Enigma subfamily of the PDZ-LIM domain protein family. To verify the interaction between Cypher and LTCC, the FLAG-tagged Cypher1c full-length or its PDZ domain (AA1–82) deletion mutant was co-expressed with the Myc-tagged LTCC cytosolic region (the last 662 amino acid residues) or a deletion mutant missing the last four residues (\(\Delta VSSL\)) in HEK293 cells (Fig. 4A). Cypher interacted with the Myc-tagged LTCC cytosolic region, whereas either the deletion mutant of the LTCC or the PDZ deletion of Cypher completely obliterated the interaction (Fig. 4A). Subsequently, the phosphorylation of LTCC cytosolic region was studied in HEK293 cells in vitro. The addition of a PKA agonist forskolin dramatically increased the phosphorylation of the LTCC (Fig. 4B). Without stimulation, the expression of Cypher slightly increased the phosphorylation of the LTCC, whereas with forskolin stimulation, the phosphorylation of the LTCC was increased further (Fig. 4B). Furthermore, the S1928A mutation completely blocked the phosphorylation of LTCC mediated by PKA (Fig. 4C). Therefore, for the cytosolic region of LTCC, Ser\(^{1928}\) was the dominant PKA catalytic site. Disrupting the interaction between LTCC and Cypher by either the \(\Delta VSSL\) deletion of the LTCC or the deletion of the Cypher PDZ domain significantly impaired the PKA-mediated phosphorylation. Our results showed that Cypher interacted with LTCC and facilitated PKA-mediated phosphorylation of the LTCC at Ser\(^{1928}\).
We further confirmed the role of Cypher in cultured cardiomyocytes utilizing virus-mediated overexpression of Cypher protein in neonatal rat cardiomyocytes. With Cypher overexpression and forskolin stimulation, phosphorylation of the LTCC was dramatically increased (Fig. 5A). We then evaluated the expression levels of the LTCC and PKA in Cypher-null mouse hearts. Cypher deletion did not change the protein levels of the catalytic subunit of PKA, the RII regulatory subunit of PKA, LTCC, nor the phosphorylated LTCC protein (Fig. 5B). Last, isoproterenol injection dramatically increased the phosphorylation of LTCC at Ser\textsuperscript{1928} in WT mouse hearts, and this was blunted in Cypher-null hearts (Fig. 5C). Similar to our in...
vitro data, PKA was tethered by Cypher to LTCC phosphorylating the LTCC at Ser1928 in cardiomyocytes.

Cypher Interacted with Ser/Thr Phosphatase CaN, Blocking Its Activity—It is a hallmark of AKAPs to bind to not only PKA, but also phosphatases or phosphodiesterases. Hence we investigated the potential interaction between Cypher and the Ser/Thr phosphatase CaN. In the mouse hearts, CaN and Cypher were in the same protein complex, as shown by co-immunoprecipitation (Fig. 6A). To confirm this interaction, tagged Cypher1c and 2c, along with the PDZ and LIM domains, were co-expressed with tagged CaN in HEK293 cells. Cypher1c and 2c interacted with CaN, whereas the N-terminal PDZ domain and C-terminal LIM domains did not (Fig. 6B). However, we discovered that different regions of Cypher2c (ΔPDZ, CCSR, and AA227–327) interacted with CaN (Fig. 6C). Deletion of the catalytic domain (AA345–354, Δcat) of CaN impaired the interaction with Cypher, whereas deletion of the AI region (AA455–477) enhanced it (Fig. 6D). Thus, Cypher interacted with CaN at the catalytic domain competing with the AI region of CaN to bind to the catalytic domain. Further, deletion of Cypher in mouse hearts did not change the protein level of CaN (Fig. 6E); however, we did detect more CaM (or CaN) protein co-immunoprecipitating with CaN (or CaM) (Fig. 6F) and increased CaN enzymatic activity (Fig. 6G). This was further evidenced by an increase in the mRNA level of modulatory calcineurin-interacting protein 1, exon 4 isoform (MCIP1.4) (32), a downstream target of calcineurin signaling, in Cypher-null hearts when compared with WT controls (Fig. 6H). Therefore, the enzymatic activity of CaN was increased in Cypher-null hearts, although the protein expression of CaN was unchanged (Fig. 6, E–H).

Cypher/ZASP Was Phosphorylated at Ser265 and Ser296 by PKA—Cypher/ZASP interacted with the RIIα regulatory subunit of PKA. Next we further analyzed whether Cypher could be phosphorylated by PKA. We searched the Cypher1c sequence for the PKA substrate consensus sequence ((K/R)(K/R)X(S/...
FIGURE 3. Cardiomyopathy-associated Cypher/ZASP mutation enhanced the PKA-Cypher interaction. A, GST-tagged human ZASP exon 4 encoded fragment and two mutants (S189L and T206I) were co-transfected with FLAG-tagged PKA RIIα (RIIα as a control) in HEK293 cells. Protein interactions were assayed by immunoprecipitation (IP), IB, immunoblotting. B, stick models of RIIα/D-Cypher Glu202-Leu215 (WT, left; T203I mutant, right) complex (RIIα D/D dimer at the back in light brown and gray, Cypher Glu202-Leu215 at the front in green) superimposed onto the RIIα D/D-AKAP-IS structure. C, surface representation of the Cypher helix (Glu202-Leu215) (WT, left; T203I mutant, right) with the highlighted hydrophobic residues (yellow) involved in PKA RIIα binding.

FIGURE 4. Cypher/ZASP tethered PKA to phosphorylate the LTCC cytosolic region at Ser1928 in vitro. A, the Cypher PDZ domain interacted with the LTCC C-terminal PDZ binding motif (VSSL). Myc-tagged Cypher or its PDZ domain deletion mutant (ΔPDZ) was co-expressed with FLAG-tagged LTCC cytosolic region (AA1510–2172) or its deletion mutant (ΔVSSL) in HEK293 cells. FLAG-tagged proteins were enriched by anti-FLAG antibody, and interacting proteins were verified by blotting with an anti-Myc antibody. IP, immunoprecipitation; IB, immunoblotting. B, FLAG-tagged LTCC cytosolic region with or without Myc-tagged Cypher was co-expressed in HEK293 cells. Forskolin (100 μM, 30 min) was used to activate PKA. LTCC protein was purified by anti-FLAG antibody, and the phosphorylation was detected by a phospho-(Ser/Thr) PKA substrate antibody. IBMX, 3-isobutyl-1-methylxanthine. C, LTCC and its mutants S1928A and C-terminal VSSL deletion (ΔVSSL) were expressed in HEK293 cells with Cypher. LTCC phosphorylation with or without the PKA agonist forskolin treatment was assessed.
is variable), and two putative PKA substrate phosphorylation sites, Ser^{265} and Ser^{296}, were identified (Fig. 7A). Ser^{265} is conserved between mouse and human, whereas Ser^{296} is unique to mouse. To begin to address the question, we first observed enhanced phosphorylation of the Cypher protein following isoproterenol treatment in the WT mouse hearts (Fig. 7B). We then analyzed whether mutants of S265A and S296A altered the phosphorylation of Cypher. Although both single mutants significantly decreased the phosphorylation of Cypher following forskolin stimulation in HEK293 cells, the double mutation S265A/S296A completely eliminated the Cypher phosphorylation (Fig. 7C).

**DISCUSSION**

Cypher/ZASP, containing a PDZ domain at its N terminus and three LIM domains at its C terminus, belongs to the PDZ-LIM domain protein family (17, 33). Cypher/ZASP is a sarcomeric protein interacting with the Z-line proteins α-actinin-2, myotilin, and calsarcin and the signaling molecule protein kinase Cs (17, 18, 34). We have demonstrated that Cypher plays essential roles in the maintenance of the sarcomere integrity especially under contraction force stress (18, 19, 30). Cypher global-null mice display postnatal death with cardiac and skeletal muscle defects. Conditional or postnatal inducible deletion of Cypher in mouse cardiomyocytes causes premature death and severe dilated cardiomyopathy. The identification of ZASP variants associated with myopathies such as dilated cardiomyopathy, left ventricular noncompaction cardiomyopathy, and myofibrillar myopathy (zaspopathy) further demonstrates the requirement of Cypher/ZASP for the normal function of striated muscle (35–37). It is a feature that the LIM domains of Enigma subfamily proteins interact with protein kinase C (38). For Cypher/ZASP, the increased affinity to PKC caused by the substitution Asp^{626} to Asn is associated with dilated cardiomyopathy (35). Moreover, deletion of Cypher or its splice isoform in mouse hearts leads to abnormalities of certain signaling pathways (18, 30). However, no direct evidence has linked Cypher/ZASP to any signal transduction pathways. Currently, our study shows that Cypher/ZASP is an AKAP. Thus, Cypher/ZASP plays not only an important structural role in sarcomeric Z-lines, but also forms a signaling center to regulate the phosphorylation of some proteins, including the LTCC and Cypher/ZASP itself. These findings are in line with the concept that cardiac Z-lines are signaling centers as well as important structural components of sarcomeres (39).

Here we provided new evidence that Cypher/ZASP is a typical type II AKAP, which tethers PKA and Ser/Thr phosphatase CaN to their specific substrates, such as LTCC, to control its function by modulating post-translational phosphorylation. The AA200–217 fragment of Cypher, forming an amphipathic helix, specifically docks to the D/D domain of PKA RIIα. Deletion of either the Cypher AA200–217 region or the RII D/D domain completely abolished the interaction (Fig. 1, F and G). The amino acid residues AA200–217 partially overlap with the
ZM motif (AA186–211), which has been assumed to be involved in the interaction with the rod domain of α-actinin (40). Interestingly, other members of the Enigma/ALP subfamily, such as ALP, which has a ZM motif (20), did not bind to PKA RIIα; however, ENH, without ZM, did bind to PKA RIIα (Fig. 1H).

FIGURE 6. Cypher directly interacted with CaN and blocked its activity. A, CaN protein was precipitated from adult mouse heart lysates, and the interaction with Cypher was verified with an anti-Cypher antibody. Rabbit IgG was used for control. IP, immunoprecipitation; IB, immunoblotting. B, GST alone or GST-tagged Cypher1c, 2c, PDZ domain (AA1–80), and LIM domains (AA547–723) were co-transfected with FLAG-tagged CaN in HEK293 cells. Interacting proteins were verified by immunoprecipitation. C, GST alone or GST-tagged Cypher1c, 2c, 2c with PDZ domain deletion (2cΔPDZ), CCSR, and 2cΔPDZPDZ was co-transfected with FLAG-tagged CaN in HEK293 cells. Interactions were verified by immunoprecipitation. D, FLAG-tagged CaN and its mutants (Δcat: Δ45–345, ΔAl: Δ455–477) were co-transfected with GST-tagged Cypher1c in HEK293 cells. Protein interactions were verified by immunoprecipitation. E, WT and Cypher-null neonatal heart lysates were probed with an anti-CaN antibody. F, CaN proteins (or CaM) were precipitated from neonatal WT, and Cypher-null hearts and interacting CaM (or CaN) were determined by probing with an anti-CaM antibody (or CaN antibody). G, CaN enzymatic activity from WT and Cypher-null neonatal heart lysates was measured (n = 4). The values were normalized to the average WT value. H, the mRNA levels of modulatory calcineurin-interacting protein 1, the exon4 isoform, MCIP1.4, in 1-day-old WT and Cypher KO mouse hearts (n = 3) measured by real-time PCR. The data are presented as the ratios of MCIP1.4 to 18 S RNA, an internal standard, and were normalized to the WT values. *, p < 0.05. Error bars indicate mean ± S.E.
phosphorylation at Ser^{1928} mediated by PKA, which is triggered by β-AR stimulation (2). However, ablation of LTCC Ser^{1928} phosphorylation in a knock-in transgenic mouse model does not affect the calcium current induced by the β-AR-CAMP-PKA signaling in mouse cardiomyocytes (45). Therefore, the actual in vivo function of Cypher/ZASP as an AKAP requires further study, including identification of the central target in the Cypher/ZASP-PKA-CaN multi-molecule signaling complex.

CaN is a Ca^{2+}/CaM dependent Ser/Thr phosphatase contributing to hypertrophic signaling in cardiomyocytes (47). Our previous work showed that deletion of the Cypher long isoforms leads to late onset dilated cardiomyopathy with overactivated CaN signaling that includes NFATc4 dephosphorylation and nuclear translocation followed by increased transcription of the NFAT downstream gene RCAN1.4 (30). The catalytic domain of CaN interacted with the middle region of Cypher/ZASP (Fig. 6, C and D). Interestingly, the deletion of the autoinhibitory region of CaN enhanced the interaction. Moreover, deletion of Cypher increased CaN-bound CaM, and therefore, the CaN enzymatic activity was increased. Several proteins have been reported to inhibit CaN activity such as calscarn (48), RCAN1.4 (49), AKAP79 (50), and FHL2 (51). It is also worthy of note that Cypher/ZASP might be a new physiologically inhibitor of CaN.

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Similarly to ENH, the PDZ domain of Cypher/ZASP binds to the LTCC at its C-terminal PDZ binding motif (31). We have demonstrated that, as an AKAP, Cypher/ZASP tethers PKA to the LTCC and phosphorylates Ser^{1928} both in vitro and in vivo. CaN, which has been reported as a phosphatase for the LTCC (41), also interacts with Cypher/ZASP. All these findings point to an additional role for Cypher/ZASP as a signaling center for regulating the Ser^{1928} phosphorylation of LTCC in cardiomyocytes. A few membrane-associated AKAPs, such as AKAP18α and AKAP79, have been reported to restrict PKA kinase activation at the LTCC (16, 42, 43). Cypher/ZASP is a dominant sarcomeric Z-line protein. The T-tubules and sarcomeric Z-lines are in close proximity within cardiomyocytes, facilitating the interaction between the sarcomeric protein Cypher/ZASP and the membrane protein LTCC, aiding in PKA regulation of LTCC channel function (44).

Our data showed that Cypher/ZASP facilitated PKA-mediated phosphorylation of LTCC at its Ser^{1928}. The S1928A substitution abolished phosphorylation of LTCC, even with forskolin treatment, which is consistent with a previous study (45). Cypher-null hearts had impaired LTCC phosphorylation at Ser^{1928} induced by isoproterenol. However, the role of Ser^{1928} phosphorylation in LTCC pump function is under debate. Ser^{1928} is a well characterized PKA/PKC/PKG phosphorylation site (46). Our data show that Ser^{1928} is the unique PKA substrate site in the cytosolic region of the LTCC (Fig. 4C). A number of in vitro studies have shown that the calcium pump function of the LTCC is regulated by modification of the
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