The novel cardiac z-disc protein CEFIP regulates cardiomyocyte hypertrophy by modulating calcineurin signaling

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The z-disc is a structural component at the lateral borders of the sarcomere and is important for mechanical stability and contractility of both cardiac and skeletal muscles. Of note, the sarcomeric z-disc also represents a nodal point in cardiomyocyte function and signaling. Mutations of numerous z-disc proteins are associated with cardiomyopathies and muscle diseases. To identify additional z-disc proteins that might contribute to cardiac disease, we employed an in silico screen for cardiac-enriched cDNAs. This screen yielded a previously uncharacterized protein named cardiac-enriched FHL2-interacting protein (CEFIP), which exhibited a heart- and skeletal muscle-specific expression profile. Importantly, CEFIP was located at the z-disc and was up-regulated in several models of cardiomyopathy. We also found that CEFIP overexpression induced the fetal gene program and cardiomyocyte hypertrophy. Yeast two-hybrid screens revealed that CEFIP interacts with the calcineurin-binding protein four and a half LIM domains 2 (FHL2). Because FHL2 binds calcineurin, a phosphatase controlling hypertrophic signaling, we examined the effects of CEFIP on the calcineurin/nuclear factor of activated T-cell (NFAT) pathway. These experiments revealed that CEFIP overexpression further enhances calcineurin-dependent hypertrophic signal transduction, and its knockdown repressed hypertrophy and calcineurin/NFAT activity. In summary, we report on a previously uncharacterized protein CEFIP that modulates calcineurin/NFAT signaling in cardiomyocytes, a finding with possible implications for the pathogenesis of cardiomyopathy.

The sarcomeric z-disc has primarily been defined as a structural component at the lateral borders of the sarcomere that is important for mechanical stability and contractility of cardiac and skeletal muscle. In the last decade, the z-disc has emerged as nodal point of cardiomyocyte signal transduction and in the pathogenesis of cardiomyopathy (1). It has become apparent that components of the z-disc like muscle lim protein (MLP)3 (2) or filamin C (3) and the costameric protein melusin (4) take part in signaling pathways or processes of mechanical stress sensing and its response. Many novel heart- and skeletal muscle-specific proteins have been discovered and suggested to play an important role in various signaling pathways, e.g. the previously described protein family of calsarcins (5) that directly interacts with the serine/threonine phosphatase calcineurin. Calsarcin-1 knock-out mice are sensitized to pathological stimuli such as pressure overload, resulting in an excessive hypertrophic response and severe cardiomyopathy (6). Overexpression of calsarcin-1 is sufficient to inhibit G_{i} agonist-induced hypertrophy and suppress calcineurin signaling in the heart (7). The calcium/calmodulin-dependent phosphatase calcineurin and its downstream targets, transcription factors of the nuclear factor of activated T-cells (NFAT) family, play essential roles in cardiomyocyte signaling (8, 9). Dephosphorylation of these transcription factors by calcineurin promotes their translocation into the nucleus and induces the pro-hypertrophic gene program. Overexpression of constitutively active calcineurin in mouse heart leads to substantial ventricular hypertrophy, massive cardiac enlargement, and heart failure (8). In contrast, mice lacking the most prevalent calcineurin isoform (CnAβ) are dramatically impaired in their ability to establish cardiac hypertrophy induced by pressure overload and angiotensin II or isoproterenol infusions (10). Similarly, the immunosuppressive agents cyclosporine A or tacrolimus (FK506) can block hypertrophic growth in cardiomyocytes by

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inhibiting calcineurin through the formation of complexes with immunophilins (11).

To identify previously unknown sarcomeric proteins, we screened expressed sequence tag (EST) databases for uncharacterized proteins predominantly found in cardiac and skeletal muscle cDNA libraries. Using a bioinformatics approach, we found an open reading frame coding for a 1435-aa protein, which we termed CEFIP (cardiac-enriched FHL2-interacting protein). This cardiac- and muscle-specific protein is dysregulated in human ischemic and dilated cardiomyopathy as well as in murine models of heart failure and hypertrophy. By performing yeast two-hybrid (Y2H) screens, we identified a well-known protein FHL2 (four and a half-LIM domains 2), which is primarily expressed in the heart, as one of the binding partners of CEFIP. FHL2 has recently been described to bind calcineurin and repress pathological cardiac growth. In contrast, we found that the overexpression of CEFIP led to dramatic induction of the hypertrophic marker genes and NFAT-dependent signaling. Taken together, we identified CEFIP as a novel z-disc protein and established its role as an activator of calcineurin-dependent hypertrophy in cardiomyocytes.

RESULTS

CEFIP is a novel cardiac- and skeletal muscle-specific protein

Numerous z-disc proteins have recently been identified as a cause for cardiomyopathies and cardiac diseases. Based on the notion that almost all of these are expressed in a tissue-restricted pattern, we performed an in silico screen for uncharacterized heart-specific expressed genes (12). Data mining and extraction mainly relied on T-STAG (Tissue-S) and Unigene EST databases. Utilizing this approach, we recently also discovered the cardiac z-disc protein Fbxl-22, which controls turnover of specific sarcomeric proteins (13). Several of the newly identified expressed sequence tags correspond to the Homo sapiens Unigene cluster Hs.585480 and the Mus musculus Unigene cluster Mm.318319 encoding C10orf71. Here, we describe a so far uncharacterized gene that we termed CEFIP (Cardiac enriched FHL2-interacting protein; human NM_001135196.1 and mouse NM_001195097.1). These NCBI reference sequences were used for designing probes containing the open reading frame. Northern blot analyses of multiple human and mouse tissues indeed showed a heart- and skeletal muscle-specific expression pattern (Fig. 1A). These results could be confirmed in quantitative real-time PCR experiments of multiple mouse tissue samples (Fig. 1B). To also analyze the expression of CEFIP at the protein level, Western blottings of several mouse tissue samples were performed. Because there was no antibody for CEFIP available, we generated a novel rabbit antiserum using a 354-aa fragment of mouse N-terminal CEFIP as immunogen. To verify the specificity of the antiserum, a multiple tissue Western blotting was probed with the novel CEFIP antibody, and another blot with antibody preincubated with the CEFIP fragment used for immunization served as control. Again, these blots revealed strong expression of CEFIP in mouse heart and skeletal muscle tissue, whereas the peptide displacement confirms the antibody’s specificity at ~250 kDa (Fig. 1C). The full open reading frame encodes a 1435-aa (human NP_001128668.1) and a 1412-aa (mouse NP_001182026.1) protein, respectively. The alignment of the CEFIP protein sequence reveals that it is highly evolutionarily conserved
CEFIP regulates cardiomyocyte hypertrophy

Figure 2. CEFIP localizes to the sarcomeric z-disc. Staining of NRVCM (A), ARVCM (B), and intact mouse heart tissue (C) with a CEFIP antibody reveals a strong signal at the sarcomeric z-disc, validated by co-localization with α-actinin (scale bar, 20 μm).

CEFIP is located at the cardiac z-disc and dysregulated in in vivo models of cardiomyopathy

To examine the subcellular localization of CEFIP, we used the newly generated antibody for immunostaining in cardiomyocytes (supplemental Fig. 2). Staining of cultivated neonatal rat cardiomyocytes with the anti-CEFIP antiserum showed a signal at the z-disc co-labeled with the well-characterized α-actinin2 antibody (Fig. 2A) (7, 14). Similarly, co-immunostaining in isolated adult rat ventricular cardiomyocytes with anti-α-actinin2 and anti-CEFIP revealed an overlapping fluorescence pattern (Fig. 2B). Subcellular immunolocalization in cryosections of intact mouse heart again confirmed z-disc localization for CEFIP (Fig. 2C).

To identify a potential patho-physiological relevance of CEFIP in vivo, we determined its expression in established models of cardiac diseases. CEFIP was significantly up-regulated in mouse models of hypertrophy like biomechanical stress due to transverse aortic constriction (TAC), calcineurin-TG mice (CnA-TG) (Fig. 3, A–D). Consistently, the expression level of CEFIP mRNA was significantly higher in samples of myocardial tissue from human patients suffering from ischemic (ICM) or dilated (DCM) cardiomyopathy compared with non-failing controls (Fig. 3E).

CEFIP modulates cardiomyocyte hypertrophy

Previously, it was shown that sarcomeric z-disc proteins play an important role in cardiomyocyte function and signaling (1). To elucidate the role of CEFIP in cardiomyocytes, we generated adenoviral constructs for overexpression or knockdown of CEFIP allowing gain and loss of function experiments. NRVCMS were infected by AdCEFIP or by a control virus over-expressing β-galactosidase (AdLacZ) and treated with the hypertrophy-inducing agent phenylephrine (PE, 100 μM). The adenoviral infection with AdCEFIP led to an overexpression of CEFIP up to 8-fold on the mRNA level and 9.7-fold on the protein level (Fig. 4, A and B), which was accompanied by an induction of members of the hypertrophic gene program such as NppA/ANF (2.36-fold), NppB/BNP (1.63-fold), Acta1 (1.9-fold), and RCAN1.4 (2-fold) (Fig. 4C). Similarly, overexpression of CEFIP significantly increased the cell-surface area (CSA), equivalent to the increased CSA due to stimulation with PE (100 μM, 24 h). Moreover, the presence of PE with AdCEFIP treatment further enhanced cardiomyocyte hypertrophy (Fig. 4, C and D). Conversely, down-regulation of CEFIP by 72% at the mRNA level and 64% at the protein level (Fig. 4, E and F), due to synthetic miRNA overexpression, revealed no differences in the expression pattern of the hypertrophic gene markers under baseline conditions, but it blunted increased expression following stimulation with PE (Fig. 4G). Interestingly, CSA is decreased in AdmiR-CEFIP-treated cardiomyocytes compared with cells treated with AdmiR-Control at baseline (Fig. 4H). Of note, AdmiR-CEFIP treatment abrogated cardiomyocyte hypertrophy induced by phenylephrine (Fig. 4H).

CEFIP is a highly dynamic protein

To establish either a potential structural or a more dynamic role for CEFIP, we studied the dynamic behavior of CEFIP with the help of FRAP. Cultured myotubes derived from immortalized mouse myoblasts were transduced with AdCEFIP-EGFP. Importantly, also in these skeletal muscle cells, EGFP-tagged CEFIP targeted to z-discs. Single z-discs in these transduced cells were bleached with a laser using a spinning disc confocal microscope (Fig. 5A and supplemental Movie S1). Analysis of CEFIP recovery showed a biphasic curve. The biexponential fit accounts for a fast and a slow phase (Fig. 5B). Because the initial fast recovery is to some extent caused by lateral diffusion of unbound EGFP-tagged protein (3), we focused on the slow halftime (1/2 slow) of CEFIP that indicates the exchange process of bound proteins with the soluble fraction. The exchange of CEFIP was extremely fast with a 1/2 slow of 32 ± 5 s, while 91 ± 3% of the protein was in the mobile fraction (Fig. 5, C and D). This half-time is comparable with that previously found for FLNC and actin in PGMS in mammalian cultured muscle cells, whereas that of α-actinin2 and especially titin is considerably slower (Fig. 5E). Therefore, CEFIP seems to belong to the group of highly dynamic z-disc proteins involved in mechanosensing, signaling, and myofibrillar remodeling (3, 16).

CEFIP directly interacts with FHL2 and induces calcineurin-dependent hypertrophy

To unravel a potential function of CEFIP in cardiomyocyte signaling, we performed Y2H experiments to identify interacting proteins. Because of auto-activation of the full-length construct, we used three different protein fragments (289–966, 966–2160, and 2160–3168 nt) as bait (Fig. 5F and Table 1). Interestingly, in screens of two fragments (966–2160 and 2160–3168 nt), we found several prey clones coding for the well-known titin and filamin-binding protein FHL2 (four and a
half-LIM domains 2). Of note, this heart-specific protein also localizes to the z-disc (17). We confirmed the interaction between HA-tagged FHL2 and Myc-tagged CEFIP via co-immunoprecipitation (Fig. 5G). Furthermore, immunostaining of adult rat ventricular cardiomyocytes confirmed co-localization of CEFIP and FHL2 at the sarcomeric z-disc (Fig. 5H). FHL2 has recently been described to directly bind to calcineurin and to repress pathological cardiac growth (17). Given this information and the pro-hypertrophic effects CEFIP exerted on cardiomyocytes, we next examined whether calcineurin signaling is affected by CEFIP. NRVCMs co-infected with a reporter adenovirus expressing luciferase under control of an NFAT-responsive promoter showed a 2.0-fold up-regulation of NFAT-luciferase activity on adenoviral overexpression of CEFIP. This effect was further enhanced in the presence of constitutively active calcineurin (Fig. 6A). Interestingly, simultaneous overexpression of CEFIP and down-regulation of FHL2 revealed a slight additive increase of luciferase activity (Fig. 6G). Conversely, knockdown of CEFIP abrogated the hypertrophic effect by the absence of FHL2 (Fig. 6H). However, overexpression of both CnA and CEFIP when FHL2 is knocked down had no significant effect compared with overexpression of CnA alone in the absence of FHL2 (Fig. 6G). Moreover, knockdown of CEFIP did not completely attenuate the activation of NFAT-luc due to the knockdown of FHL2 in the presence of CnA (Fig. 6H). Thus, it is unlikely that FHL2 and CEFIP function in a linear pathway. Instead, we hypothesize that CEFIP plays an independent role to fine-tune calcineurin/NFAT signaling in cardiomyocytes (Fig. 7).

Discussion

Here, we introduce CEFIP as a novel striated muscle-enriched sarcomeric z-disc protein. This previously uncharacterized protein appears to play an important role in car-
CEFIP regulates cardiomyocyte hypertrophy

A

CEFIP mRNA expression [n-fold]

Control CEFIP

B

CEFIP Tubulin

Control CEFIP

C

mRNA expression [n-fold]

NppA NppB Acta1 RCAN1.4

Control Veh. Control PE CEFIP Veh. CEFIP PE

D

relative cell surface area [n-fold]

Vehicle PE Vehicle PE

E

CEFIP mRNA expression [n-fold]

miR-Ctr. miR-CEFIP

F

CEFIP Tubulin

miR-Ctr. miR-CEFIP

G

mRNA expression [n-fold]

NppA NppB Acta1 RCAN1.4

miR-Ctr. Veh. miR-Ctr. PE miR-Ctr. Veh. miR-CEFIP PE

H

relative cell surface area [n-fold]

Vehicle PE Vehicle PE
CEFIP regulates cardiomyocyte hypertrophy via modulation of calcineurin-dependent signaling.

**CEFIP is a novel z-disc protein**

Recently, the perception of the z-disc has experienced a transformation from the traditionally, merely mechanical role into a nodal point in cardiomyocyte signaling (1) and a “hotspot” of cardiomyopathies. Several z-disc components have been discovered to participate in the processes of mechanical stress sensing and signal transduction (3, 18, 19). Because of its relatively large size, one might assume that CEFIP plays a role as a structural component of the z-disc. However, FRAP experiments revealed unforeseen dynamics and mobility of CEFIP compared with other sarcomeric proteins in mammalian (3, 18, 19) and avian (20, 21) muscle cells implying a role in signaling rather than a mere structural function within the myofibril.

**CEFIP interacts with FHL2 and regulates hypertrophic signaling**

Because the function of CEFIP was largely unknown, we first applied a yeast two-hybrid screen using three independent protein fragments of CEFIP as baits in order to identify possible protein interaction partners. We identified FHL2 as a binding partner of CEFIP and could verify this interaction by co-immunoprecipitation and co-localization. Interestingly, at least two independent binding sites for FHL2 must exist on CEFIP, as two non-overlapping baits revealed this new interaction partner. FHL2, like CEFIP, is localized at the z-disc and highly enriched in the heart (22). FHL2 interacts with a multitude of proteins that are localized to the cell membrane, cytoplasm, and sarcomere as well as to the nucleus (23). Calcineurin, one of the key players in cardiac hypertrophy (8), ranks among these proteins (17). Yet the role of FHL2 in cardiac hypertrophy is still controversial. The absence of FHL2 in vivo does not produce a pathological phenotype at baseline. Furthermore, FHL2-deficient mice develop hypertrophy to the same extent as wild-type mice following TAC (24). In contrast, the hypertrophic response is exaggerated in FHL2 knock-out mice when stimulated with the β-adrenergic receptor agonist isoproterenol (25). While under hypertrophic conditions, the absence of FHL2 seems to produce none or maladaptive effects, and the induction of FHL2 appears to be beneficial. Overexpression of FHL2 blunts hypertrophy upon stimulation with phenylephrine or MEK1 (26). Consistently, the inhibition of hypertrophy in ROCK2-deficient mice upon stimulation with angiotensin II is mediated by enhanced expression of FHL2 (27). Interestingly, an increase or reduction of CEFIP expression is associated with opposite effects in terms of cardiomyocyte hypertrophy compared with FHL2.

**CEFIP affects calcineurin signaling**

Our results indicate that overexpression of CEFIP is sufficient to induce cardiomyocyte hypertrophy. Moreover, the hypertrophic effects of phenylephrine are exaggerated by CEFIP overexpression. Conversely, knockdown of CEFIP blunts hypertrophy induced by phenylephrine. 

\( G_{q}/G_{11} \) agonists like angiotensin II or phenylephrine mediate hypertrophy through several molecular pathways (28), including the calcineurin–NFAT cascade (29). In line with this notion, knockdown of CEFIP is sufficient to inhibit hypertrophy induced by calcineurin, which is also an interaction partner of FHL2 (17). Activation of calcineurin leads to dephosphorylation of NFAT transcription factors and their translocation to the nucleus, where they initiate the induction of hypertrophy-associated genes (30). Overexpression of CEFIP increases NFAT–luciferase–reporter activity and even further enhances the effect of constitutively active calcineurin. Similar results have been obtained by knockdown of FHL2 before (17) and could be reproduced by us. However, combination of CEFIP overexpression with FHL2 knockdown during overexpression of constitutively active calcineurin was not able to further augment promoter activity, which might be due to a ceiling effect. Of note, the increase of calcineurin-dependent NFAT promoter activity caused by the absence FHL2 is abolished by knockdown of CEFIP. Taken together, the interaction of FHL2 with calcineurin as well as with CEFIP establishes a complex of these three proteins at the z-disc, where FHL2 and CEFIP generate opposing effects on calcineurin/NFAT signaling (Fig. 7).

**CEFIP in cardiac disease**

We also provide data that CEFIP is induced in human failing hearts due to ischemic or dilated cardiomyopathy. We found a similar expression pattern in different animal models with heart failure. Again, FHL2 expression seems to be changed in the opposite direction: FHL2 is down-regulated in human non-ischemic heart failure (31). Moreover, a missense mutation of FHL2 was identified in a patient with familial dilated cardiomyopathy (32). FHL2 is suppressed in hypertrophic cardiomyopathy as well (33).

Integrating the heart- and skeletal muscle-specific expression of CEFIP and its induction in models of cardiac disease, we hypothesize that CEFIP acts as a mediator in cardiac disease. It may be part of the maladaptive hypertrophic response of cardiomyocytes to biomechanical stress that accompanies valvular or ischemic heart disease. Moreover, one may speculate that...
gain of function mutations of CEFIP trigger cardiac hypertrophy. To clarify the role of CEFIP in heart disease mutation analyses of CEFIP in patients with dilated or hypertrophic cardiomyopathy might be helpful.

Because we could clearly show that overexpression of CEFIP is sufficient to induce hypertrophy by modulating the calcineurin pathway, CEFIP represents a potential target for a therapeutic intervention such as siRNA attempts. Because of its heart and skeletal muscle specificity, negative or unwanted side effects might be minimized.

In summary, we describe here a novel cardiac- and skeletal muscle-specific protein termed CEFIP, which functions as a regulator of calcineurin-dependent hypertrophy. On the one hand, increased CEFIP expression led to an induction of car-

Figure 5. CEFIP is a highly dynamic protein that interacts with FHL2. A–E, FRAP analysis of CEFIP dynamics in z-discs of cultured myotubes expressing EGFP-tagged CEFIP. A, CEFIP is targeted to z-discs. ROI for bleaching was limited to a single z-disc, and fluorescence recovery was followed over time. Time series of fluorescence micrographs are shown before bleaching (pre-bleaching), immediately after bleaching (post-bleaching), and 5 and 20 s after bleaching and after maximum fluorescence recovery. B, typical recovery profile of CEFIP in z-discs. C, slow halftime of CEFIP in z-disc of 32.2 s (median) is shown as box plot of all experiments (n = 8). D, estimation of CEFIP mobility in z-discs indicates that 91% of the protein is in the mobile fraction (n = 8). E, overview of exchange half-lives (t1/2) and mobile fractions (Mf) of known heart related proteins. F, because of auto-activation of the full-length CEFIP, three yeast two-hybrid screens were performed using different protein fragments of CEFIP as bait. In two screens, several clones encoding FHL2 were found. G, FHL2 was confirmed as a partner of full-length CEFIP via co-immunoprecipitation (IP) from HEK293 cells expressing full-length Myc-tagged CEFIP and HA-tagged FHL2. H, co-immunostaining of adult rat ventricular cardiomyocytes confirms co-localization of CEFIP and FHL2 in z-disc. Scale bars, 5 μm (A), 10 μm (H). WB, Western blotting.
diomyocyte hypertrophy; on the other hand, the absence of CEFIP repressed pharmacological agonist- and calcineurin-induced hypertrophy. Additional in vivo experiments will help to clarify the role of CEFIP in cardiac-signaling pathways and in the pathogenesis of cardiomyopathy.

Experimental procedures

**Northern blot analysis**

Multiple tissue Northern blots (BioChain) containing human or mouse poly(A) RNA were hybridized with [32P]dCTP-labeled (Rediprime II Random Prime Labeling System, Amer sham Biosciences) cDNA probes corresponding to the ORF of human or mouse CEFIP at 65 °C overnight. Washing was repeated with 2× SSC, 0.1% SDS at 80 °C for 24–168 h. Autoradiography was conducted at −80 °C for 24–168 h with an intensifying screen.

**Isolation and culture of NRVCMs**

NRVCMs were isolated according to a previously published protocol (19). Hearts from 1- to 2-day-old Wistar rats (Charles River Laboratories) were excised and minced in ADS buffer (120 mM NaCl, 20 mM HEPES, 8 mM NaH2PO4, 6 mM glucose, 5 mM KCl, 0.8 mM MgSO4, pH 7.4). An enzymatic solution containing collagenase type II (0.5 mg/ml, Worthington) and pan creatin (0.6 mg/ml, Sigma) in sterile ADS buffer was used to digest the tissue. A Percoll (GE Healthcare) gradient centrifuga tion step was applied to remove contaminating fibroblasts from cardiomyocytes. NRVCMs were resuspended and cultured in Dulbecco’s modified Eagle’s medium (DMEM) containing 10% FCS, 1% penicillin/streptomycin, and 1% l-glutamine (PAA) for 24 h. After 24 h, cells were infected with adenoviruses in serum-free medium for a further 24 h and afterward stimulated with phenylephrine for 48 h.

**Isolation and culture of adult rat ventricular cardiomyocytes (ARVCMs)**

ARVCMs were isolated using collagenase and plated on laminin-coated dishes as described (34). ARVCMs were cultured in a HEPES-modified medium 199 (M199, Sigma S7528, supplemented with 5 mM taurine, 5 mM carnitine, 5 mM creatine, 5 mM N-mercaptopropionyl glycine, 0.1 μM insulin, 10,000 units/ml penicillin, and 10 mg/ml streptomycin, pH 7.25).

**Immunofluorescence microscopy**

NRVCMs, ARVCMs, and mouse tissue cryosections were fixed with 4% paraformaldehyde in PBS for 10 min at room temperature, permeabilized, and blocked with 2.5% BSA and 0.1% Triton X-100 in PBS for 1 h at room temperature. The slides were incubated with a monoclonal mouse antibody for α-actinin 2 (1:400, Sigma) together with polyclonal CEFIP (rabbit 1:100) antibody at 4 °C overnight. The CEFIP antibody was generated by immunizing rabbits with a purified, bacterially expressed recombinant protein representing the first 354 amino acids of mouse CEFIP (custom immunization by Biogenes, Berlin, Germany). Its specificity was confirmed by Western blotting. Secondary antibodies against mouse immunoglobulins conjugated with Cy3 (Dianova) and rabbit immunoglobulins conjugated with FITC (Dianova) were used at a dilution of 1:500 for 1 h at room temperature. Simultaneously, nuclei were labeled with DAPI (4’,6’-diamidino-2-phenylindole, Vector Laboratories), and after serial washing with PBS, the slides were mounted with FluorPreserve reagent (Merck).

**Table 1**

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<td>TAO kinase 3</td>
<td>51347</td>
<td>STE20 like kinase</td>
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</table>
Confocal images were taken with a Zeiss LSM800 laser-scanning microscope with a Plan-Apochromat ×40/1.4 oil differential interference contrast (UV) visible-IR objective at room temperature. Cropped regions from the overviews are newly acquired images. Cell-surface areas were captured and analyzed on BZ-9000-E HS all-in-one fluorescence microscope (Keyence) at ×60 magnification (Plan-Apo oil immersion ×60, NA 1.4; Nikon) or ×20 magnification (CFI Plan-Apo ×20, NA 0.75; Nikon).

**RNA isolation and real-time PCR**

Total RNA from NRVCMS was isolated using QIAzol lysis reagent (Qiagen), according to the manufacturer’s protocol,

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**Figure 6. CEFIP interacting partner FHL2 shows no additive effect on the NFAT-luciferase activity.**

A, for luciferase reporter assays NRVCMS were co-infected with adenoviruses expressing NFAT reporter-based firefly luciferase, Renilla luciferase, constitutively active calcineurin, and CEFIP or β-galactosidase (LacZ as control). Overexpression of CEFIP increases the NFAT-luciferase activity 2.02 ± 0.1-fold (n = 9, p < 0.01). B, overexpression of constitutively active calcineurin produces induction of hypertrophic marker genes (NppA, NppB, and Rcan1.4). Overexpression of CEFIP on top of calcineurin exaggerates this effect (NppA: 1.66 ± 0.1-fold, n = 9, p < 0.01; NppB: 1.77 ± 0.2-fold, n = 9, p < 0.05; Rcan1.4: 3.1 ± 0.6-fold, n = 9, p < 0.05). C, overexpression of calcineurin results in increased cell-surface area. CEFIP enhances hypertrophic growth induced by calcineurin in NRVCMS (1.2 ± 0.02, n > 2800, p < 0.001). D, infection of NRVCMS with different quantities of synthetic miRNA down-regulating CEFIP (12.5, 25, 50 m.o.i.) reduces dose-dependently the NFAT-luciferase activity down to 30.96 ± 4.5% (n = 9, p < 0.001). E, adenoviral knockdown of CEFIP blunts the induction of NppA, NppB, and Rcan1.4 caused by constitutively active calcineurin as shown by qPCR. F, consistently, down-regulation of CEFIP inhibits hypertrophic growth induced by calcineurin in neonatal rat ventricular cardiomyocytes. G, luciferase expression driven by constitutively active calcineurin is amplified in NRVCMS infected with an adenovirus coding for CEFIP (9.14 ± 0.9-fold, n = 14, p < 0.001). This induction is not further increased by the addition of siRNA knockdown of FHL2. 

H, induction of the calcineurin driven luciferase activity by absence of FHL2 could not be blunt by simultaneous knockdown of CEFIP. Statistical significance was calculated by two-way ANOVA. Error bars show means ± S.E. *p < 0.05; **p < 0.01; ***p < 0.001.
and was DNase I-digested. After purification, 1 μg of RNA was reverse-transcribed into cDNA using the Superscript III first strand cDNA synthesis kit (Life Technologies, Inc.). For qRT-PCR, the EXPRESS SYBR GreenER Reagent (Life Technologies, Inc.) or iQ Multiplex Powermix (Bio-Rad) was used in the CFX96 real time cycler (Bio-Rad).

Cloning of human and mouse CEFIP

The complete open reading frame of CEFIP was amplified from heart cDNA using the following gene-specific primers: mouse CEFIP_F, 5'-GCT GGC ACC ATG CAG GGA AAC AAA AAA TGC AC-3', and mouse CEFIP ms_R, 5'–GCT GGG TCG CCT CAA GAA ACA CCC TCT GTG GCA AAG-3'; human CEFIP_F, 5'-GCT GGC ACC ATG ATG CAA GGA AAT AAG AAG TGC ACA GAC G-3', and human CEFIP ms_R, 5'-GCT GGG TCG CCT CAA GAA ATG CCT TCT GTG GCA AAG-3'. The PCR product was recombined into a pDONR221 gateway vector using the Gateway Technology (Life Technologies, Inc.). For protein expression, CEFIP was shuttled into an expression plasmid or entry vector to design constructs coding for N-terminal fusion tags (HA, MYC, and V5). An adenovirus encoding the full-length CEFIP was generated with the pAd/CMV/V5-DEST Gateway vector kit following the manufacturer's instructions. As a control, a β-galactosidase-V5 encoding adenovirus was used for overexpression experiments.

Cloning of synthetic CEFIP knockdown miRNA

Oligonucleotides against CEFIP were designed using Invitrogen's BLOCK-iT RNAi Designer (miR-CEFIP_TOP, TGC TGA GCA GAA GTT GAC ACC ACT TTG TTT TGG CCA CTG ACT GAC AAA GTG GTC AAC TTC TGC T and GCC CAA AAC AAA GTG GTG TCA ACT TCT GCT C) and cloned into a pcDNA 6.2-GWmiR vector according to the manufacturer’s protocol. This construct was recombined into pDONR221 entry vector and shuttled into pAd/CMV/V5 adenoviral expression vector. For knockdown experiments, the pcDNA6.2 GWmiR plasmid, which is predicted not to target any known mammalian gene, served as a control.

Fluorescence recovery after photobleaching (FRAP) and data analysis

C2C12 myotubes were transduced with CEFIP-EGFP and kept at 37 °C and 5% CO₂. FRAP experiments were performed essentially as described (3, 16) using a Cell Observer Spinning Disk Confocal Microscope (Carl Zeiss, Jena, Germany) equipped with an external 473-nm laser coupled via a scanner (UGA-40, Rapp OptoElectronic, Hamburg, Germany) and evaluated using Zen 2012 software (Carl Zeiss). For FRAP analysis, eight independent experiments were performed. The region of interest (ROI) for bleaching was limited to a single z-disc. For each cell, 1–3 ROIs were chosen and bleached using a 473-nm laser (100 milliwatts) with 100% intensity (Rapp OptoElectronic) for a pulse time of 1 ms with eight iterations. Images were taken before and immediately after bleaching, and fluorescence recovery was monitored for 230 s with an interval time of 0.1–1 s until the signal was fully recovered. Raw data were transformed into normalized FRAP curves as described previously (3, 16).

Yeast two-hybrid library assays

Automated Y2H screens were performed as described (35). Human cDNA libraries from human heart and skeletal muscle (Clontech) as well as a library of individual cloned full-length open reading frames from cDNAs of 5000 different genes (DKFZ Heidelberg) were screened to a minimal coverage of 5 million clones per library. As full-length human CEFIP showed auto-activation, the screen was performed using three nucleotide (nt) fragments of CEFIP (289–966, 966–2160, and 2160–3168 nt). To mate yeast strains harboring the bait protein and
CEFIP regulates cardiomyocyte hypertrophy

the prey library, exponentially growing cultures of an absorbance (600 nm) of 1 were combined, pelleted by centrifugation, and resuspended in an equal volume of YPDA medium containing 20% PEG 6000. Mating mixes were incubated at 30 °C with gentle agitation for exactly 3 h before washing and resuspending cells in selective medium. Briefly, clones displaying differential growth on selective plates lacking histidine, leucine, and tryptophan were picked and grown in selective medium lacking leucine and tryptophan, and plasmid DNA was isolated and subsequently electroporated into DH10B *Escherichia coli* (Gibco via Invitrogen, Karlsruhe, Germany). The obtained clones were sequenced and retransformed with the CEFIP construct to confirm the interaction. Interaction pairs were chosen to be real pairs if out of two independent bacterial preparations from the same yeast clone the sequenced clone is identical, or where the sequence of two preys that interacted with the same bait is highly related.

**Luciferase reporter gene assays**

NRVCMS were infected with adenoviruses coding for either CEFIP (25 m.o.i.) for overexpression experiments or synthetic miRNA against CEFIP (100 m.o.i.) for knockdown experiments. An adenovirus encoding LacZ as well as an adenovirus encoding unspecific miRNA was used as controls. AdNFATluc carrying miRNA against CEFIP (100 m.o.i.) for knockdown experiments. AdRen-luc carrying Renilla luciferase were co-infected. 72 h after infection, cells were harvested in passive lysis buffer (Promega). The luciferase assay was performed using the dual-luciferase reporter assay (Promega) according to the manufacturer’s protocol on the infinite m200 PRO system (Tecan). The experiments were performed in quadruplicate and repeated at least three times.

**Co-immunoprecipitation**

HEK293 cells were transfected with plasmids encoding the potential CEFIP interacting proteins (N-terminal fusions) by using JetPEI (PolyPlus Transfection) according to the manufacturer’s instructions. Cells were harvested 48 h post-transfection with ice-cold ELB lysis buffer (50 mM HEPES, pH 7.0, 250 mM NaCl, 1% Nonidet P-40, 5 mM EDTA, Protease Inhibitor Mixture (Roche Applied Science, catalog no. 1836170)) containing Phosphatase Inhibitor Mixture 2 and 3 (Sigma, catalog no. P5726; P0044). After incubation on ice for 30 min, lysates were centrifuged for 5 min at 16,200 × g at 4 °C to remove cellular debris. A total of 3000 µg of cleared protein extract was incubated with anti-HA-agarose (Sigma, catalog no. A2095) for 2 h at 4 °C with moderate agitation. After four cycles of washing with lysis buffer, protein complexes were eluted in 1× bead volume SDS sample buffer. For Western blot experiments, the beads were separated by 2 min of centrifugation at 16,200 × g at room temperature, and the supernatant containing the protein fraction was analyzed by immunoblotting.

**Western blotting**

To determine specificity of CEFIP expression, protein extracts from NRVCMS or tissue samples were isolated and proceeded in Lysis-Buffer (20 mM Tris, pH 7.5, 12.5% (v/v) glycerol, 10 mM dithiothreitol, 500 mM NaCl, 1% (v/v) Nonidet P-40 (Sigma), 10 mM DTT, as well as protease inhibitor mixture (Roche Applied Science) and phosphatase inhibitor mixture 2 and 3 (Sigma)). Protein concentration was measured by Bradford protein assay kit according to the manufacturer’s guidelines (Bio-Rad). Samples were resolved by SDS-PAGE, transferred onto a PVDF membrane (GE Healthcare), and incubated with respective target-specific primary antibody. Subsequently, the membrane was incubated in secondary HRP-coupled antibody followed by visualization using a chemiluminescence kit (ECL, GE Healthcare) and detection with a FluorChemQ imaging system (Alpha Innotech).

**Statistical analyses**

All results are the means ± S.E. Statistical analyses were performed using two-tailed Student’s *t* test, one-way or two-way ANOVA followed by Student-Newman-Keuls post hoc tests when appropriate. *p* values < 0.05 were considered statistically significant.

**Author contributions**—F. D. conducted experiments, analyzed and interpreted the data, and wrote the manuscript. C. K. designed the study and interpreted data. C. R. performed the experiments and analyzed the data. S. H. performed experiments. S. S. provided the human samples. D. O. F., P. F. M. v. d. V., J. B., and S. M. performed and interpreted the FRAP experiments and generated the CEFIP-antibody. N. F. devised the strategy and designed the study. All authors critically revised the manuscript.

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**References**


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