The complement C3-complement factor D-C3a receptor signalling axis regulates cardiac remodelling in right ventricular failure

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Failure of the right ventricle plays a critical role in any type of heart failure. However, the mechanism remains unclear, and there is no specific therapy. Here, we show that the right ventricle predominantly expresses alternative complement pathway-related genes, including Cfd and C3aR. Complement 3 (C3)-knockout attenuates right ventricular dysfunction and fibrosis in a mouse model of right ventricular failure. C3a is produced from C3 by the C3 convertase complex, which includes the essential component complement factor D (Cfd). Cfd-knockout mice also show attenuation of right ventricular failure. Moreover, the plasma concentration of CFD correlates with the severity of right ventricular failure in patients with chronic right ventricular failure. A C3a receptor (C3aR) antagonist dramatically improves right ventricular dysfunction in mice. In summary, we demonstrate the crucial role of the C3-Cfd-C3aR axis in right ventricular failure and highlight potential therapeutic targets for right ventricular failure.

Cardiovascular disease is the leading cause of death worldwide1. Substantial efforts have been made in studying the pathophysiology of heart failure due to its high prevalence and poor prognosis. Most of the previous studies have focused on left ventricular (LV) failure since the left ventricle plays a central role in supplying blood to the systemic circulation against strong resistance1. Therefore, several medical therapies for heart failure have been developed, such as beta-blockers, angiotensin-converting enzyme inhibitors, and mineralocorticoid receptor antagonists. Right ventricular (RV) failure has not been a central issue because the right ventricle pumps blood solely to the pulmonary artery against weak resistance1. However, accumulating evidence indicates that RV failure significantly contributes to the pathophysiology of various types of heart failure1-3. RV failure is one of the most important predictors of symptoms and prognosis in patients with pure right-sided heart disease3. Additionally, RV failure is a strong independent risk factor for survival in patients with LV failure5-8.
However, drugs developed for treating LV failure are not effective against RV failure, and no effective therapies specifically targeting RV failure exist. Hence, we aimed to understand the RV failure pathophysiology and develop an effective treatment for RV failure. The right ventricle differs from the left ventricle in many ways, including its structure, function, and developmental origin. In terms of cellular properties, cardiomyocytes in the right ventricle are different from those in the left ventricle. In humans, certain diseases predominantly affect the right ventricle. These findings indicate that the right ventricle has unique characteristics, and hence, there exist therapeutic targets specific to RV failure. Here, we focused on the right ventricle to dissect the unique molecular signature of RV failure and develop a specific therapy. We believe our findings could pave way for development of RV failure-specific therapeutics.

Results

Complement system activation in the right ventricle

To understand the molecular signature of the right ventricle, we first screened for genes specifically expressed in the right ventricle. We separated the murine heart tissue into the left ventricle, right ventricle, and ventricular septum and performed transcriptome analysis. Each cardiac tissue showed differential gene expression patterns for cardiac-specific genes (Supplementary Fig. 1a–f). Global gene expression analysis indicated that each cardiac region showed similar but different gene expression patterns (Fig. 1a). Ingenuity pathway analysis (IPA) of the differentially expressed genes between the right ventricle and left ventricle or between the right ventricle and ventricular septum showed that genes involved in the complement system were significantly enriched in the right ventricle, represented by Cfd, C3, and C3ar1 (Fig. 1b, c). Importantly, Cfd, C3, and C3ar1 are also highly expressed in human RV (Fig. 1d–f). During complement activation, the C3 protein is cleaved into C3a and C3b by C3 convertase complex factors, including complement factor D (Cfd), and C3ar is the receptor of C3a. Therefore, we surmised that the complement system might play an important role in RV functions. To investigate whether the complement system was altered under pathological conditions, we generated an RV failure mouse model induced by pulmonary artery constriction (PAC). Interestingly, PAC also increased the expression of Cfd and C3ar1 in the right ventricle (Fig. 1g–i). We then confirmed RV dysfunction and upregulation of cardiac failure and fibrotic markers specifically in the right ventricle (Fig. 1j, k; Supplementary Fig. 1g–j). Since C3a is hardly detectable in histological analysis and is a cleaved fragment of C3, C3d is used as a marker of C3 activation and is covalently fixed to tissues. C3d was detected in the right ventricle only after PAC (Fig. 1l). These data suggest that the complement system was activated in the right ventricle, and it might contribute to the pathophysiology of RV failure.

C3 is a critical factor for RV failure development

C3 is a central component of the complement system, and its activation stimulates several downstream pathways. To explore the possibility that depletion of C3 modulated the progression of RV failure, we performed PAC in wild type (WT) and C3−/− mice. Interestingly, RV systolic dysfunction and dilatation induced by PAC were attenuated in C3−/− mice compared with WT mice (Fig. 1m; Supplementary Fig. 2a–d). These data suggest that C3 contributes to the development of heart failure specifically in the right ventricle.

The liver-derived C3 plays a critical role in RV failure development

Although plasma C3 is mainly produced in the liver, C3 is also produced in other tissues where it functions locally. To clarify the origin of C3, which contributed to the development of RV failure, we generated a C3 floxed mouse model by conventional homologous recombination (Supplementary Fig. 3a, b). We then created the cardiac- or liver-specific C3 knockout mice by crossing C3 floxed mice with α-myosin heavy chain promoter-driven Cre (αmHC-Cre) or albumin promoter-driven Cre (Alb-Cre) mice, respectively, and performed PAC surgery. Intriguingly, PAC induced RV dysfunction and dilatation in the cardiac-specific C3 knockout mice, but these changes were attenuated in the liver-specific C3 knockout mice (Fig. 2a, b; Supplementary Fig. 3c–e). RV fibrosis was attenuated only in the liver-specific C3 knockout mice, but left ventricle was not affected (Fig. 2c–f; Supplementary Fig. 3f–i). Consistently, cardiac failure and fibrotic markers were significantly upregulated in the cardiac-specific C3 knockout mice compared with the liver-specific C3 knockout mice (Fig. 2g–j). These data suggest that the liver-derived C3 plays a crucial role in the development of RV failure.

Cfd/CFD is a critical factor for RV failure

In the complement activation cascade, cleavage of C3 by C3 convertase generates the active molecules C3a (anaphylatoxin) and C3b (opsin). Cfd is required for the generation of the C3 convertase complex and is essential for producing C3a. Since Cfd was enriched in the right ventricle, we next focused on the C3-Cfd signalling axis in the pathogenesis of RV failure. We performed PAC in Cfd−/− mice, which showed no apparent gross abnormalities at baseline. Similar to C3−/− mice, PAC-induced RV dysfunction and dilatation were significantly suppressed in Cfd−/− mice, but left ventricle was not affected (Fig. 3a, b; Supplementary Fig. 4a–c). RV fibrosis, cardiac failure, and fibrotic marker gene expression were also significantly suppressed in Cfd−/− mice compared with WT mice (Fig. 3c–g; Supplementary Fig. 4d, e).

Based on the findings that the C3-Cfd signalling axis contributed to the development of RV failure in mice, we decided to examine whether the complement system was also involved in human RV failure pathogenesis. We examined the plasma concentrations of C3 and CFD in patients with chronic RV failure and compared these with the severity of the disease represented by the mean pulmonary artery (PA) pressure and plasma B-type natriuretic peptide (BNP) level. Plasma C3 concentration did not correlate with plasma BNP levels or mean PA pressure (Supplementary Fig. 4f, g). However, plasma CFD concentration was significantly correlated with plasma BNP levels and mean PA pressure (Fig. 3h, i). These data suggest that CFD was correlated with the severity of human RV failure.

C3a directly regulates cardiac gene expression

Since C3ar1 was highly expressed in the right ventricle, we focused on the C3a-C3ar1 signalling axis in the heart. We examined whether C3a regulated cardiac-specific gene expression in cultured rat cardiomyocytes. Several types of mitogen-activated protein kinases (MAPKs) play a central role in cardiac remodelling and heart failure. The addition of
C3a activated extracellular signal-regulated kinase (ERK) in a time-dependent manner and weakly activated p38, but it did not activate c-Jun N-terminal kinase (JNK) (Fig. 4a). To examine whether C3a-dependent ERK phosphorylation was mediated by C3aR, we used siRNA to knockdown C3ar1. As expected, C3ar1 knockdown blocked C3a-dependent ERK activation (Fig. 4b). Next, to understand the role of C3a in cardiomyocytes, we performed a global gene expression analysis of cardiomyocytes cultured in the presence or absence of C3a stimulation (Fig. 4c; Supplementary Fig. 5a). IPA of genes enriched in the presence of C3a stimulation showed that the heart failure- and inflammation-related pathways were involved (Fig. 4d). We then searched for genes that showed changes in expression similar to that observed in our in vivo analyses. Among the genes enriched under C3a stimulation (Supplementary Fig. 5a), the expression of Ccl5 and Cebpa was significantly upregulated in WT mice with PAC and suppressed in C3−/− mice with PAC (Supplementary Fig. 5b, c). Therefore, these genes could be the downstream targets of the C3a-C3aR signalling axis in RV failure.
C3aR blockade ameliorates the development of RV failure after PAC
From a clinical point of view, we investigated the applicability of the C3a-C3aR pathway in treating RV failure. We used a non-peptide C3aR antagonist, SB290157, to selectively block C3aR signalling. SB290157 administration significantly suppressed the development of RV dysfunction after PAC in WT mice (Fig. 4e–f, Supplementary Fig. 6a–c). Furthermore, increases in the weights of the liver, lung, and heart, which are signs of heart failure, were suppressed by SB290157 treatment (Supplementary Fig. 6d–f). Fibrosis and elevation of cardiac fibrosis and fibrotic markers in the right ventricle were also attenuated in mice with PAC by SB290157 treatment, but left ventricle was not affected (Fig. 4g–j, Supplementary Fig. 6g–k).

Given that arrhythmia could be a fatal complication of RV failure, we investigated whether SB290157 affected the incidence of arrhythmia in mice with PAC. RV free wall tachypacing easily induced ventricular tachyarrhythmia in mice with PAC, but not in sham-operated mice. Interestingly, C3−/− and WT mice with SB290157 treatment significantly suppressed the induction of ventricular tachyarrhythmia and reduced the duration of ventricular tachyarrhythmia in mice with PAC (Fig. 4k–m). Calcium homeostasis plays a pivotal role in cardiomyocyte excitation-contraction coupling, and its impairment leads to RV failure and arrhythmic events. In isolated RV cardiomyocytes, time to peak Ca2+ transient was prolonged by PAC in WT mice, and its prolongation was suppressed in C3KO mice (Fig. 4n–o). The frequency of spontaneous Ca2+ waves was increased by PAC in WT RV cardiomyocytes, and its frequency was attenuated in C3KO RV cardiomyocytes (Fig. 4p–q). Therefore, these data suggest that abnormal Ca2+ dynamics are manifested in RV failure, and C3KO ameliorates its aberrant Ca2+ dynamics.

Discussions
RV functioning integrates preload, afterload, contractility, configuration, size, pericardial constraint, and interaction with the left ventricle. Dysregulation of each of these factors induces RV failure. Its underlying causes include pulmonary vascular disease, par enchymal lung disease, RV infarction, congenital heart diseases, and LV failure. Although therapies for some specific diseases have been developed, specific therapies for RV failure have not been developed. To develop a drug for RV failure, it is important to understand its pathogenesis by murine RV failure models. Constriction of the pulmonary artery could induce less systemic or toxic effects, but a pulmonary vascular disease model of chronic pressure overload-induced RV failure.

Understanding innate immunity helps us to understand the role of the complement system in sterile inflammation. Recent data show the possible involvement of the complement system in LV failure. The complement system is activated in patients with LV failure, and complement factors are associated with clinical outcomes. Low levels of plasma C3 can be a predictive marker of early death in patients with LV failure and increased levels of complement factor B and Bb are associated with mortality in patients with LV failure. Animal model studies also uncovered the role of the complement system in LV failure. Pulmonary arterial hypertension is one of the important diseases to induce RV failure and is induced by hypoxia in experimental animals. The complement system activation was observed in the perivascular lesion in hypoxic lung, and would be involved in the pathology of pulmonary hypertension. These basic and clinical studies focused solely on LV failure and lung, and not on RV failure. Upon activation of the complement system, the central complement factor C3 is cleaved. Cfd is a rate-limiting enzyme in the alternative pathway and controls subsequent processes. In the right ventricle, Cfd is predominantly expressed. C3 is highly abundant, but C3a is quickly converted by carboxypeptidase N into C3a-desArg and C3a-desArg cannot bind to C3aR. Therefore, we speculated that the C3a-C3aR signalling axis was only involved in RV failure. C3a-mediated signalling has been described as proinflammatory or anti-inflammatory; thus, these effects are context-specific. It remains unclear why complement-related genes are highly expressed in the right ventricle. The complement system fundamentally prepares the body for infections by a lytic cascade activation, but is a complex innate immune surveillance system, playing a role in host homeostasis, inflammation, and in the defense against pathogens. The right ventricle is in the venous return circulation system and is possibly at a risk of pathogen and foreign material intrusion. Therefore, basal expression of complement-related genes may play a role in surveillance and protection under physiological conditions. Nonetheless, in the current study, it remains unclear whether the upstream complement pathway could be activated and as to how this pathway is maintained in RV failure. The C3aR antagonist rescued the pathological phenotype, and we speculated that the downstream pathway, the membrane attack complex, would not play a role in RV failure. The C3aR antagonist rescued the pathological phenotype, and we speculated that the downstream pathway, the membrane attack complex, would not play a role in RV failure. Therefore, it would be interesting for future studies to determine whether the downstream signalling could be activated in RV failure. The C3aR antagonist, SB290157 is widely used to explore the role of C3aR in animal experiments, but may have other effects, such as a partial C5aR agonist. In drug study, we should take off target effect into consideration.

In summary, we showed that the right ventricle specifically expressed Cfd and C3aR, and their expression was exaggerated in PAC-induced RV failure. Whole body C3 deletion, Cfd deletion, and the liver-specific C3 deletion ameliorated PAC-induced RV failure in mice. In patients with RV failure, the CFD concentration was significantly correlated with the severity of RV failure. C3a directly regulated the expression of several genes through C3aR in cardiomyocytes, and the C3aR antagonist ameliorated PAC-induced RV failure in mice. Currently, there is no specific biomarker for patients with RV failure, and hence, we speculate that CFD could be a disease marker for RV failure in humans. Anaphylatoxins have recently been explored as therapeutic targets for several diseases. Here, we showed the crucial role of the C3-Cfd-C3aR signalling axis in RV failure and highlighted the potential therapeutic targets for RV failure, which has no pharmacologic options...
at present. It would be interesting to develop C3-, C3a-, or CFD-targeted drugs for RV failure treatment.

**Methods**

**Animal**

This study conformed to the Guide for the Care and Use of Laboratory Animals published by the US National Institutes of Health (Publication no. 85-23, revised 1996), and the study protocol was approved by the Institutional Animal Care and Use Committee at the Keio University School of Medicine. To generate C3-flx mice, a targeting vector was constructed as follows: a 1.1 kb DNA fragment carrying exons 24 and 25 of the C3 gene was amplified by polymerase chain reaction (PCR) and inserted between the KpnI sites of the middle entry clone (pDME-1). This clone consisted of a DNA fragment of pgk promoter-driven

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5
Neo-poly(A) flanked by two frt sites, and two loxP sequences located 123 bp upstream of exon 24 and 108 bp downstream of exon 25, respectively. The 5.8 kb upstream and 7.2 kb downstream homologous genomic DNA fragments were subcloned into the 5′ entry clone (pDSU2-2) and 3′ entry clone (pD3DE-2), respectively. For targeting vector assembly, the three entry clones were recombined into a destination vector plasmid (pDEST-ΔT; containing a cytomegalovirus enhancer/chicken actin (CAG) promoter-driven diptheria toxin gene) using MultiSite Gateway Technology (Thermo Fisher Scientific). Homologous recombinant embryonic stem (ES) clones were identified by Southern blot and PCR analyses. ES cell culture and chimeric mice generation were performed as previously described35. C3/MHC-Cre mice (C57BL/6 N background), α-miosin heavy chain promoter-driven Cre mice (αMHC-Cre) and albumin promoter-driven Cre mice (Albumin-Cre; C57BL/6 background) were purchased from Jackson Laboratory. αMHC-Cre mice and Albumin-Cre mice were backcrossed to the C57BL/6 N background. Cardiac-specific C3-MHC knockout mice were generated by crossing C3/MHC-Cre mice with αMHC-Cre mice. Liver-specific C3 mice were generated by crossing C3/MHC-Cre mice with Albumin-Cre mice. C3MHC mice (C57BL/6 N background) were generated using the CRISPR/Cas9 system, as described previously36. In all genetically modified mice, genotypes were confirmed by PCR analysis (please see Supplementary Table 1 for list of primer sequences).

Establishment of the pulmonary artery constriction (PAC) mouse model
PAC surgery was performed on 8-week-old C57BL6 mice as previously described35. The surgery was performed using a ventilator to acquire passive respiration, and anaesthesia was induced with 2% isoflurane. The main trunk of the pulmonary artery was constricted using a 25-gauge blunt needle as the calibrator. The pulmonary artery blood flow was detected by Doppler echocardiogram analysis 2 weeks after surgery. The sham operation followed the same procedure, except that the pulmonary artery was not constricted. Mice were examined for the subsequent analyses 2–4 weeks after surgery.

Establishment of the transverse aortic constriction (TAC) mouse model
TAC surgery was performed on 8-week-old mice, as previously described35. The surgery was performed using a ventilator to acquire passive respiration, and anaesthesia was induced with 2% isoflurane. The aorta was constricted using a 27-gauge blunt needle as the calibrator. The innominate artery and left common carotid artery blood flows were detected by Doppler echocardiogram analysis 1 week after surgery. The sham operation followed the same procedure, except that the aorta was not constricted. Mice were examined for the subsequent analyses 4 weeks after surgery.

Echocardiography
Mice were anaesthetised using isoflurane, and echocardiography was performed using ultrasonography (Vevo 2100 system, Visual Sonics, Toronto, Canada) with a 30 MHz probe. Stable images were obtained in the M-mode, B-mode, and Doppler Mode. The LV inner dimension and fractional shortening were measured using the M-mode. The RV inner dimension and fractional shortening were measured using the B-mode. The pulmonary arterial pressure gradient was obtained using the Doppler Mode. Heart rate did not differ significantly among the experimental groups during the echocardiographic assessments. All analyses were performed in a blinded manner with respect to the mice genotype.

In vivo electrophysiological study (EPS)
An in vivo EPS was performed 2–4 weeks after PAC or sham operation. Mice were anaesthetised in a mixture of 0.3 mg/kg medetomidine, 4.0 mg/kg midazolam, and 5.0 mg/kg butorphanol. The heart rate was maintained at 400–500 bpm during the EPS. Programmed electrical stimulations (burst pacing and extrastimulus pacing) were delivered from the right ventricle using 1.1-Fr electrophysiology catheters (EPR-900, Millar, Houston, TX, USA), and electrocardiograms were recorded by electrodes placed on the extremities.

In vivo hemodynamic study
LV and RV function were assessed by pressure catheter in vivo37. Mice were anaesthetised using isoflurane. The right carotid vein was exposed, and a 1.4-Fr pressure catheter (SPR 839; Millar Instruments, Houston, TX) was inserted into the RV while recording in digital form (MPVS PL3508 PowerLab 8/35, ADInstruments) at the acquisition rate of 2 kHz for analysis (LabChart 8 pro, ADInstruments). Next, the left carotid artery was exposed, and a 1.4-Fr pressure catheter was inserted into the LV while recording LV pressure. All values were averaged over five consecutive cardiac cycles during stable phase of the respiratory cycle.

Administration of the C3a receptor (C3aR) antagonist SB290157
C3aR antagonist SB290157 (Sigma-Aldrich) was administered (10 mg/kg body weight/day) using an Alzet micro-osmotic pump (model 2002). Alzet micro-osmotic pumps were implanted subcutaneously into the intrascapular region of mice on the day following PAC operation. During the pump implantation, the mice were anaesthetised with isoflurane.

Transcriptome analysis
Mice were deeply anaesthetised and sacrificed by decapitation. Sternotomy was performed, cold phosphate-buffered saline (PBS) was perfused into the right and left ventricles, and the hearts were extirpated and separated into the right ventricle, left ventricle, and ventricular septum. The separated cardiac tissues were immediately frozen in liquid nitrogen. Frozen heart tissues were broken into powder using Cryopress (CP-100W, Microtec-Nichon Co. Ltd, Funabashi, Japan). These powders were dissolved in TRDzol reagent (Innogenit), and total RNA was extracted. Similarly, cultured neonatal rat ventricular cardiomyocytes (NRVCs) were washed with cold PBS and immediately dissolved in TRDzol reagent. Transcriptome analysis was performed using SurePrint G3 Mouse Microarray 8X60 K ver.2.0 (Agilent Technologies Inc., Santa Clara, CA, USA) and SurePrint G3 Rat GE 8 × 60K ver.2.0 (Agilent Technologies Inc.). The chips were scanned using Agilent Scanner G2505C (Agilent Technologies Inc.). The data
Fig. 3 | Complement factor D (Cfd) plays an important role in mice with pulmonary artery constriction (PAC)-induced right ventricular (RV) dysfunction and patients with RV failure. a, b Measured values obtained from the echocardiogram in wild type (WT) sham, WT PAC mice, and Cfd knockout (Cfd KO) PAC mice (n = 4, 5, 3, p < 0.001, p = 0.0017). The right ventricle (RV) contractile function (right ventricular fractional shortening [RV FS]) and RV size (right ventricular end-diastolic diameter [RVDd]) were evaluated. Data are presented as mean ± standard deviation (SD). c Representative images of hematoxylin-eosin (HE) staining and Azan staining of the heart in Cfd KO PAC mice. LV, left ventricle (n = 4). d, e Quantified fibrotic area of the RV and LV in WT PAC and Cfd KO PAC model mice (n = 4). Data are presented as mean ± SD. f, g qRT-PCR analysis of the expression of heart failure markers (Nppb and Myh7) in the LV, ventricular septum, and RV of WT sham, WT PAC, and Cfd KO PAC mice (n = 3–5, p = 0.0135, p = 0.0005). Data are presented as mean ± standard error of the mean (SEM). In qRT-PCR analysis, expression of target genes was normalised to that of Gapdh. Significance was assessed using a two-tailed unpaired Student’s t-test. *p < 0.05; **p < 0.01. h Scatter plots showing the correlation between the CFD concentration and B-type natriuretic peptide (BNP) concentration in the overall cohort (n = 129; mean age = 66.5 ± 15.1 years; 69.8% women). Spearman correlation coefficient and two-tailed p-value are shown. Linear regression line (blue line) with 95% confidence intervals (grey area) is represented. i Scatter plots showing the correlation between the CFD concentration and mean pulmonary artery (PA) pressure in the overall cohort (n = 133; mean age = 66.1 ± 15.3 years; 70.7% women). Spearman correlation coefficient and two-tailed p-value are shown. Linear regression line with 95% confidence intervals is represented.
were processed and analysed using the GeneSpring software (v.14.1.1, Agilent Technologies Inc.). Statistical significance was assessed using analysis of variance (ANOVA) ($p < 0.05$) and corrected for multiple-group comparisons using Tukey’s HSD. Significant differentially expressed genes were captured, and the expression patterns of these genes were analysed for pathway analysis using the Ingenuity Pathway Analysis software (Ingenuity Systems, www.ingenuity.com).

Histology
Mice were euthanized by cervical dislocation. The heart was immediately perfused with PBS and fixed with 10% neutral buffered formalin. The tissues were then embedded in paraffin. Sections were cut to a thickness of 4 μm and stained with hematoxylin and eosin. The connective tissue was visualised by Azan staining. For immunostaining, the hearts were perfused with PBS and fixed with 4% paraformaldehyde.
Tissues were placed in 10 and 20% sucrose in PBS until they sank (1–2 h) and then in 30% sucrose in PBS overnight. For the detection of C3d, the samples were incubated with mouse monoclonal anti-C3d (3d29, Creative BioLabs) primary antibody for 1 h at 37 degrees. Antibody binding and then in 30% sucrose in PBS overnight. For the detection of C3d, the sections were then counterstained with tetramethyl rhodamine isothiocyanate (TMRITC, Molecular Probes). The fluorescence images were captured on a confocal microscope (LSM 880, Carl Zeiss). The confocal images were then merged with the bright-field images to generate the merged images.

**Isolation of adult cardiomyocytes**

The process of adult cardiomyocyte isolation in mice was basically performed as described previously. After 1 h of serum starvation (1% fetal bovine serum), NRVCs were stimulated with C3a recombinant protein (8085-03-025, R&D SYSTEMS). siRNAs were transfected into cells using Lipofectamine3000 (Invitrogen) according to the manufacturer’s protocol. The siRNA for C3ar1 (s136363) was purchased from Thermo Fisher Scientific. Control siRNA and scrambled siRNA (4390846) were purchased from Life Technologies.

**Real-time quantitative reverse transcription PCR (qRT-PCR) analysis**

In brief, total RNA was purified using the RNeasy Mini Kit (Qiagen). RNA samples were treated with dDNA Remover (Toyobo) to remove genomic DNA contamination. One microgram of DNase-treated RNA was used for first-strand complementary DNA (cDNA) synthesis using the ReverTra Ace qPCR RT Kit (Toyobo) and oligo dT20 primers. qPCR was performed using Fast SYBR Green Master Mix (Thermo Fisher Scientific). The primer sequences are listed in Supplementary Table 2.

**Western blot analysis**

Briefly, the NRVCs were lysed using the ULTRARIPA kit A solution (BioDynamics Laboratory Inc., Tokyo, Japan) containing 1% protease inhibitor (P8340, Sigma-Aldrich) and 1% phosphatase inhibitor (160-24371, Wako Pure Chemical Industries Ltd.). The cell lysates were centrifuged at 12,000 × g, and the protein concentration in the supernatant was determined using the BCA Protein Assay Kit (TaKaRa Bioc Inc., Japan). Total proteins were resolved in 15% sodium dodecyl sulfate polyacrylamide gels under reducing conditions and electrophoretically transferred onto polyvinylidene difluoride membranes using the iBlot Dry Blotting system (Thermo Fisher Scientific). The membranes were incubated with Blocking One (Nacalai Tesque, Kyoto, Japan) for 30 min to block nonspecific binding sites. The polyclonal antibody was then incubated overnight with the primary antibodies diluted in Hikari Signal Enhancer Solution (Nacalai Tesque) at 4 degrees. Then, the membranes were incubated with horseradish peroxidase-conjugated anti-rabbit IgG (NA934, GE Healthcare) secondary antibody for 1 h at room temperature. Labelled proteins were visualised using an enhanced chemiluminescence kit (Nacalai Tesque) according to the manufacturer’s instructions. Primary antibodies related to MAP kinases were rabbit anti-p44/42 MAPK (4695, Cell Signalling Technology), rabbit anti-phospho-p44/42 MAPK (4495, Cell Signalling Technology), and rabbit anti-phospho-p38 (9211, Cell Signalling Technology).

**Measurement of Ca2+ Imaging**

Briefly, isolated cardiomyocytes were loaded with Tyrode solution containing 1 μM Fluo-4AM (F14217, Thermo Fisher Scientific) and stained for 20 min. The loaded cardiomyocytes were then resuspended in normal Tyrode solution containing 1 mM CaCl2 and kept at 37 degrees. Electric field stimulation (DFS-007, DIA Medical System Co) was applied to loaded cardiomyocytes. During this process, Ca2+ transients, Ca2+ waves, and Ca2+ sparks were captured on a confocal microscope (LSM 880, Carl Zeiss). The parameters of Fmax/F0, Time to Peak, and RT50 were calculated from fluorescence intensity in the cardiomyocytes along a 100 μm line. The time at which the fluorescence intensity decreased to 50% of Fmax was defined as Ca2+ wave. The time at which the fluorescence intensity decreased to 40% of Fmax was defined as Ca2+ spark. The Ca2+ wave transient was measured as the change in fluorescence intensity of Fluo4 during 5 Hz electric field stimulation for approximately 5 s. Ca2+ waves and Ca2+ sparks were analyzed as the change in fluorescence intensity emitted from isolated cardiomyocytes during 15 s after field stimulation. Ca2+ waves were counted as fluorescence intensity in 90% of the surface area of cardiomyocytes. The Ca2+ sparks were counted as a localized Ca2+ spark in the cardiomyocytes along a 100 μm line. The parameters of Fmax/F0, Time to Peak, and RT50 were calculated from the Ca2+ transients. Fmax/F0 was calculated by defining F0 as the fluorescence intensity of Fluo4 at rest before field stimulation and Fmax as the maximum fluorescence intensity of Fluo4 during field stimulation. Time to Peak was calculated as the difference between the time of the first electric field stimulation and the time of the first Fmax. RT50 was calculated as the difference between the time at Fmax and the time at which the fluorescence intensity decreased to 50% of Fmax.

**Human sample correction**

We retrospectively evaluated patients with chronic thromboembolic pulmonary hypertension (CTEPH) and pulmonary arterial hypertension (PAH) from April 2016 to November 2020 according to our criteria. The patients were divided into two groups: CTEPH patients and PAH patients. The CTEPH patients were defined as patients with pulmonary hypertension due to chronic thromboembolic disease. The PAH patients were defined as patients with primary pulmonary hypertension. The characteristics of the patients are listed in Table 1.
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Author contributions
S.I. and S.Y. designed the experiments. S.I., H.H., H.Y., D.K., Y.A., T.N., M.M., J.K., T.K., M.Y., K.S., A.K., M.L., M.S., G.Y., C.M., T.S., T.Y., Y.S., T.H., M.K., Y.K., K.Suzuki, K.I., M.A., M.O., H.T., M.Y., K.Sakimura, M.Y., and S.Y performed the experiments. M.M., Y.S., T.H., M.K., and T.K collected the human samples. K.I. and H.Y. conducted the mouse physiological study. H.Y. conducted Ca2+ imaging study. H.T. and M.Y. generated the Cdf knockout mice. K.Suzuki, M.A., K.Sakimura and M.Y. generated the conditional C3 knockout mice. Y.F. and K.F. supervised the study. S.I. and S.Y. wrote the manuscript. All authors read and approved the final manuscript.

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