BASIC SCIENCE

Cardiac mesenchymal stromal cells are a source of adipocytes in arrhythmogenic cardiomyopathy

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Aim
Arrhythmogenic cardiomyopathy (ACM) is a genetic disorder mainly due to mutations in desmosomal genes, characterized by progressive fibro-adipose replacement of the myocardium, arrhythmias, and sudden death. It is still unclear which cell type is responsible for fibro-adipose substitution and which molecular mechanisms lead to this structural change. Cardiac mesenchymal stromal cells (C-MSC) are the most abundant cells in the heart, with propensity to differentiate into several cell types, including adipocytes, and their role in ACM is unknown. The aim of the present study was to investigate whether C-MSC contributed to excess adipocytes in patients with ACM.

Methods and results
We found that, in ACM patients’ explanted heart sections, cells actively differentiating into adipocytes are of mesenchymal origin. Therefore, we isolated C-MSC from endomyocardial biopsies of ACM and from not affected by arrhythmogenic cardiomyopathy (NON-ACM) (control) patients. We found that both ACM and control C-MSC express desmosomal genes, with ACM C-MSC showing lower expression of plakophilin (PKP2) protein vs. controls. Arrhythmogenic cardiomyopathy C-MSC cultured in adipogenic medium accumulated more lipid droplets than controls. Accordingly, the expression of adipogenic genes was higher in ACM vs. NON-ACM C-MSC, while expression of cell cycle and anti-adipogenic genes was lower. Both lipid accumulation and transcription reprogramming were dependent on PKP2 deficiency.

Conclusions
Cardiac mesenchymal stromal cells contribute to the adipogenic substitution observed in ACM patients’ hearts. Moreover, C-MSC from ACM patients recapitulate the features of ACM adipogenesis, representing a novel, scalable, patient-specific in vitro tool for future mechanistic studies.

Keywords
Arrhythmogenic cardiomyopathy • Mesenchymal stromal cells • Adipogenesis • Fibrofatty substitution • Plakophilin2 • Plakoglobin

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**Translational perspective**

Fibro-adipose substitution has a double detrimental effect on the myocardium in arrhythmogenic cardiomyopathy (ACM), worsening arrhythmogenesis by creating a non-conductive substrate, and causing ventricular dysfunction leading to heart failure. Notably, to-date no etiological therapy is available. This work introduces, for the first time, the stromal cardiac compartment as a key player in ACM ventricular adipose substitution: we demonstrated that cardiac human mesenchymal stromal cells undergo adipogenic differentiation both in ACM explanted hearts and in culture through a PKP2-dependent mechanism. Cardiac mesenchymal stromal cells constitute a suitable cellular platform for future mechanistic studies and a potential target for future therapies.

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**Introduction**

Arrhythmogenic cardiomyopathy (ACM) is a genetic disease in which the ventricular myocardium, predominantly the right ventricle, is progressively substituted by fibro-adipose tissue. Arrhythmogenic cardiomyopathy is associated with severe ventricular arrhythmias, sudden death, and progressive heart failure.

Fibro-adiposis is the hallmark of the disease; it typically starts from the epicardium at the ‘triangle of dysplasia’ and progresses toward the endocardium. Arrhythmogenic cardiomyopathy has a double pathologic phenotype: arrhythmias and ventricular dysfunction. The pathogenic mechanisms for both phenotypes are unclear. Although a primary electrical cause of arrhythmias involving sodium channels and connexins has been proposed, fibro-fatty substitution may worsen the condition of patients’ by creating non-conducting tissue. Moreover, progressive adipocyte deposition leads to loss of cardiac contractility and ultimately heart failure.

The disease usually has an autosomal dominant transmission with variable penetrance and expressivity. Rare recessive forms of ACM occur in conjunction with skin manifestations. Approximately 50% of symptomatc individuals harbour a mutation in one of the genes of the five major members of the cardiac desmosome: DSP (desmoplakin), PG (plakoglobin), PKP2 (plakophilin2), DSG2 (desmoglein2), and DSC2 (desmocollin2). Desmosomes are complex cellular junctions found mainly in epidermal cells and in cardiac myocytes. Non-desmosomal genes TGFβ3, TMEM43, LMNA, DES, TTN, PLN, CTNNB1, RYR2, and a number of other loci have also been associated with ACM.

It has been suggested that defective desmosomes cause stretch-induced cardiomyocyte death and impair intracellular signaling that determines cell fate. As a consequence, a cell differentiation hypothesis has been formulated to explain the fibro-adipose replacement, involving the inhibition of canonical WNT/β-catenin signaling, an important regulator of myogenesis vs. adipogenesis.

A key question in ACM pathogenesis is the cellular origin of excess adipocytes. The signal for enhanced fibro-adipogenesis is likely to originate from cells that express desmosomes. Mature cardiomyocytes, second heart field cardiovascular progenitor cells (c-KIT/SCA1), or sub-epicardial progenitor cells are the most extensively studied candidates. Mesenchymal stromal cells (MSC) are supportive cells found in many tissues. They are multipotent and, under appropriate stimuli, can differentiate into osteoblasts, chondroblasts, and adipocytes. Mesenchymal stromal cells are characterized by the expression of mesenchymal surface antigens such as CD44, CD105, CD29, and CD90. In our laboratory, a population of cardiac mesenchymal stromal cells (C-MSC) isolated from adult human auricles have been characterized. In a normal adult heart, cardiomyocytes only represent 30% of the total cell number. The remaining 70% consists of cells other than cardiomyocytes, among which C-MSC are the vast majority. Cardiac mesenchymal stromal cells play a critical role in maintaining normal cardiac function, as well as in cardiac remodelling during pathological conditions. Mesenchymal stromal cells switch between osteoblast and adipocyte differentiation has been reported as a causative mechanism of osteoporosis.

The present study addresses, for the first time, the role of C-MSC in ACM adipose substitution in human hearts. We provide evidence that in ACM hearts cells differentiating into adipocytes express mesenchymal markers and C-MSC isolated from human ACM hearts express desmosomal genes and are more prone to accumulate fat (lipogenesis) and specifically differentiate into adipocytes compared with C-MSC from control hearts.

**Methods**

Detailed/additional methods are presented in Supplementary material online.

**Ethical statement**

This study complies with the declaration of Helsinki. Arrhythmogenic cardiomyopathy (ACM) patients’ samples use for research was approved by Centro Cardiologico Monzino and La Sapienza University Ethic Committees. Samples from patients not affected by arrhythmogenic cardiomyopathy (NON-ACM) were obtained from cadaveric donors from Fondazione Banca dei Tessuti di Treviso.

**Cardiac mesenchymal stromal cells culture and adipogenic differentiation**

Cardiac mesenchymal stromal cells were isolated, cultured, and characterized as previously reported. Adipogenic differentiation was obtained using adipogenic medium for 72 h or 1 week. Fat accumulation was tested by Oil Red O (ORO) or Nile Red staining; adipogenesis was tested by anti-Perilipin1 (PLIN1) antibody staining.

**Gene expression analysis**

qRT-PCR was performed in duplicate using 10 ng cDNA obtained after total RNA extraction with TRizol, treatment with Dnase I, and reverse transcription with Superscript III according to manufacturer’s instructions. Primer sequences are reported in Supplementary material online, Table S4.

**Protein expression analysis**

Western blots were performed by standard procedures, after total protein extraction by Laemmli buffer lysis, target proteins were detected with
| Sample type | ACM patient ID | Sex | Type/age of first manifestation | Dysfunction and structural alterations | Tissue characterization of wall | Repolarization ab. | Depolarization/conduction ab. | Arrhythmias | Family history | Mutations in ACM-associated genes |
|-------------|----------------|-----|---------------------------------|----------------------------------------|-------------------------------|-------------------|-----------------------------|----------------------------|----------------|----------------------------------|
| EH → FFPE   |                |     |                                 |                                        |                               |                   |                             |                           |                |                                   |
| H1          | F 42           |     | VT/34                           | Major                                   | Major                         | Major             | Major                       | Major          | Neg.             | n.a.                             |
| H2          | F 37           |     | Sincope, sustained VT/33        | Major                                   | Major                         | Neg.              | Minor                       | Minor          | Major             | DSP: c.4198C > T                  |
| H3          | M 53           |     | VT/11                           | Major                                   | n.a.                          | Neg.              | Minor                       | Major          | Neg.             | n.a.                             |
| Biopsy → FFPE |              |     |                                 |                                        |                               |                   |                             |                           |                |                                   |
| B1          | F 37           |     | Palpitations, polymorphic PVC/37| Minor                                   | Minor                         | Neg.              | Minor                       | Minor          | Neg.             | None*                             |
| B2          | M 48           |     | Sustained VT/34                 | Minor                                   | Major                         | Neg.              | Minor                       | Minor          | Neg.             | None                             |
| B3          | M 26           |     | ECG– inverted T waves/26        | Neg.                                    | Major                         | Neg.              | Minor                       | Minor          | Neg.             | None                             |
| Biopsy → C-MSC |            |     |                                 |                                        |                               |                   |                             |                           |                |                                   |
| B4          | M 54           |     | Cardiac arrest/50               | Neg.                                    | Minor                         | Major             | Neg.                        | Major          | Major             | PKP2: c.2013delC                  |
| B5          | M 44           |     | Cardiac arrest/42               | Minor                                   | n.a.                          | Major             | Neg.                        | Major          | Major             | PKP2: c.1881delC not prev. described |
| B6          | M 41           |     | PVC/32                          | Major                                    | Not conclusive                | Major             | Neg.                        | Minor          | Major             | PKP2: c.1643delG                  |
| B7          | M 51           |     | Sustained VT/50                 | Minor                                   | n.a.                          | Major             | Neg.                        | Major          | Neg.             | None                             |
| B8          | M 57           |     | Polymorphic PVC/56              | Neg.                                    | Minor                         | Minor             | Neg.                        | Minor          | Neg.             | None                             |
| B9          | F 38           |     | ECG- inverted T waves/37        | Neg.                                    | Neg.                          | Major             | Neg.                        | Minor          | Major             | PKP2: c.2013delC                  |

FFPE, formalin-fixed, paraffin-embedded tissue section; Echo, 2D-echocardiography; CMR, cardiac magnetic resonance; AG, angiography; n.a., not available; PVC, premature ventricular contractions; VT, ventricular tachycardia; EH, explanted heart; neg., negative; ab., abnormalities.

*Only PKP2 was examined.
primary antibodies PKP2, PG, DSP1/2, DSG2, DSC2, PPARγ, PLIN1, FABP4, and GAPDH (see Supplementary material online, Table S3).

Plakophilin2 overexpression
Cardiac mesenchymal stromal cells were transfected with control GFP or wild-type PKP2-GFP vector using the Neon Transfection System.

Cardiac mesenchymal stromal cells and heart section immunofluorescence
Cells were fixed with 4% paraformaldehyde, paraffin was removed from embedded sections and antigen unmasking was performed. Primary and secondary antibodies staining was carried out in 2% serum.

Statistical analysis
Descriptive variables are reported as mean ± standard deviation. Results were analysed by two-tailed Student’s t-test, Pearson’s correlation test or Fisher’s exact test. Benjamini Hochberg correction was applied when appropriate. Results were considered statistically significant at $P < 0.05$.

Results
Cells differentiating into adipocytes in arrhythmogenic cardiomyopathy human hearts are of mesenchymal origin
To understand the cellular origin of excess adipocytes in ACM hearts, we performed double staining with PLIN1 antibody and mesenchymal markers CD29 and CD105 on serial slices of three ACM explanted hearts (H1–3, Table 1; Figure 1) and three autopsy control hearts (see Supplementary material online, Table S1). PLIN1 is a protein, specifically expressed in adipocytes, surrounding lipid droplets. We observed contour staining of large lipid drops of mature adipocytes, as well as small droplets which eventually fuse during maturation of immature adipocytes. We found that all of these differentiating cells express the mesenchymal markers CD29 and CD105 (Figure 2A). We did not detect PLIN1 staining in cells expressing α-sarcomeric actin (αSARC; Figure 2B), which suggests that mature cardiomyocytes are not the source of adipogenesis. Further, since the involvement of cardiac progenitor cells was suggested, we looked for immature adipocytes expressing the c-KIT marker. We detected occasional PLIN1 and c-KIT
double-positive differentiating cells (see Supplementary material online, Figure S1). CD29 and CD105-positive differentiating cells were also found in left ventricle (LV) samples of patients H1 and H2, showing biventricular ACM, whereas only few cells were observed in control hearts (data not shown). Moreover, to verify that pre-adipocytes are of mesenchymal origin even at early phases of the disease, we obtained paraffin sections (FFPE) of biopsies from three patients with mild forms of ACM (B1–3, Table 1). We found fewer pre-adipocytes than in explanted hearts; however, all stained positive for mesenchymal markers (see Supplementary material online, Figure S2A and B). In addition, we did not detect any PLIN1-positive cells co-staining for c-KIT (see Supplementary material online, Figure S2C).

Cardiac mesenchymal stromal cells can be obtained from arrhythmogenic cardiomyopathy hearts and studied in vitro

To verify whether the MSC cardiac population may represent the source of excess adipocytes in ACM hearts, C-MSC were isolated from right ventricular endomyocardial biopsy samples from six ACM patients (B4–9, Table 1) and from five control subjects’ right ventricular autopsy samples (see Supplementary material online, Table S1). The biopsy procedure was performed for diagnostic purpose (Figure 3E and F) in patients with electrocardiogram (ECG; Figure 3A) and cardiac magnetic resonance (CMR; Figure 3B and Supplementary material online, Videos A and B) findings suggestive of ACM. Biopitic samples for C-MSC isolation (Figure 3D) were acquired in the area adjacent to the electro-anatomical scars (Figure 3C), without evidence of adipose substitution.

Cardiac mesenchymal stromal cells express desmosomal genes and proteins

Cardiomyocytes or their progenitors have been previously studied in ACM as the only cardiac cell types expressing desmosome genes. As shown by gene expression analysis in Figure 4A, we demonstrated that desmosomal genes PKP2, JUP, DSG2, DSP, and DSC2 are detectable in isolated C-MSC, and expressed at similar levels in ACM and control cells. Whole ventricle tissue samples were examined as positive controls. This result raised the possibility that C-MSC may be directly affected by ACM-causing mutations. Western blot analysis confirmed that PKP2, PG, DSP, and DSC2 proteins were expressed in ACM and control C-MSC. Plakophilin2 expression was lower in ACM C-MSC (Figure 4B).

Arrhythmogenic cardiomyopathy cardiac mesenchymal stromal cells show more propensity than controls to differentiate into adipocytes

When cultured in adipogenic medium (72 h and 1 week), ACM C-MSC accumulated lipids earlier and in significantly higher amounts than NON-ACM control cells (Figure 5). Indeed, after 72 h in adipogenic medium, lipid content, quantified by ORO staining, showed statistically different values between NON-ACM and ACM cells.
The expression of genes involved in adipogenesis or specific for adipocytes was then examined. We found that the enhanced expression of PPARγ, ADIPOQ, FABP4, and PLIN1 after 72 h in adipogenic medium was significantly more pronounced in ACM than in NON-ACM C-MSC (Figure 6A). This result was confirmed by western blot for PPARγ, FABP4, and PLIN1 (Figure 6C). As shown in Figure 6B, a positive correlation exists between ADIPOQ and PLIN1 expression and lipid accumulation analysed by ORO staining ($r = 0.874$ and $r = 0.884$, respectively).

In contrast, the anti-adipogenic gene CTGF$^{34}$ and the cell-cycle regulator CCND1 were expressed at significantly lower levels in ACM samples than in NON-ACM controls (Figure 6A).

To test the involvement of the WNT pathway, we treated ACM C-MSC with 6-bromoindirubin-3′-oxime (BIO), a drug that inhibits GSK3β$^{35}$. C-MSC cultured with BIO showed a partial (31.25%) reduction in accumulated lipid droplets ($P = 0.002$) and a 92.24% reduction in ADIPOQ expression ($P = 0.045$; see Supplementary material online, Figure S4). Notably, PG showed a preferential nuclear localization in ACM C-MSC and a predominant localization in the cytoplasm and at cell–cell junctions in NON-ACM C-MSC (see Supplementary material online, Figure S5).

**Lipid accumulation and transcriptional rearrangements in arrhythmogenic cardiomyopathy cardiac mesenchymal stromal cells are dependent on PKP2 deficiency**

In order to establish whether the diminished expression of PKP2 plays a causal role in lipid accumulation, ACM C-MSC were transfected either with a plasmid encoding PKP2, or with an empty control vector. PKP2 overexpression was confirmed both at gene and protein levels (Figure 7A). After 72 h in adipogenic medium, cells were examined...
for lipid accumulation and expression of adipogenic genes. Oil Red O staining showed a significant decrease in the number of PKP2-overexpressing cells containing lipid droplets when compared with mock-transfected controls (Figure 7C; 21.33% ± 4.65 vs. 32.57% ± 4.71, P = 0.047). Moreover, PKP2-overexpressing C-MSC showed a 98.10% and a 29.80% decrease in ADIPOQ (P = 0.034) and PPARγ (P = 0.032) expression, respectively (Figure 7B). Finally, we confirmed that lipid accumulation is dependent on PKP2 deficiency by using a specific siRNA during adipogenic induction. Indeed, silencing of PKP2 (see Supplementary material online, Figure S6A) was accompanied by a significant increase in lipid accumulation (15.87 ± 6.99 vs. 38.66 ± 16.91; P = 0.009; see Supplementary material online, Figure S6B).

**Arrhythmogenic cardiomyopathy cardiac mesenchymal stromal cells undergo adipogenesis**

To verify that the accumulation of lipid droplets is associated with C-MSC differentiation to adipocytes and not to mere lipogenesis, we analysed the expression of PLIN1 by qRT-PCR, western blot (Figure 6A and C) and immunofluorescence (Figure 8A). Indeed, ACM C-MSC accumulated lipid surrounded by adipocyte-specific PLIN1 protein. ACM C-MSC cultured in adipogenic medium stained with anti-PLIN1 antibody are phenotypically very similar to the adipogenic differentiating cells detected in ACM patients’ hearts (Figures 2 and 8B). Furthermore, ACM C-MSC maintained the mesenchymal marker CD29 during adipogenesis in vitro (see Supplementary material online, Figure S7).

**Discussion**

To date, ACM has been considered a disease confined to contractile cells even if there is no conclusive experimental evidence to support this hypothesis. To our knowledge, this work is the first demonstration that desmosomal genes are expressed in C-MSC, besides cardiomyocytes, and epithelial cells.12,35,36 Cardiomyocytes have been shown to accumulate lipid in different models, but each of them...
exhibit shortcomings. Indeed, induced pluripotent stem cells (iPSC)-derived cardiomyocytes are a promising patient-specific tool to study ACM as they accumulate lipids.22,23,37 Ongoing studies might overcome the lack of terminal differentiation in cardiomyocytes derived from iPSC, and the presence of residual partially reprogrammed cells.38 Evidence of desmin and vimentin expression in cells differentiating into adipocytes in heart tissue of a suspected ACM patient has been reported21; however, these proteins are not expressed exclusively in cardiomyocytes and adipocytes. Another study claimed that neutral lipids accumulated in ACM right ventricular myocardial tissue38; despite this, there was no evidence of myocardial cell differentiation into adipocytes. Finally, transgenic murine models of ACM, expressing different disease-causing mutations under the αMHC promoter,6,16 showed the accumulation of adipocytes in the heart, but trans-differentiation of cardiomyocytes into adipocytes was not investigated. In human ACM specimens, we found neither PLIN1-positive lipid droplets accumulating in cardiomyocytes nor αSARC staining in immature adipocytes: this rules out a direct trans-differentiation of cardiomyocytes into adipocytes. Nevertheless, we cannot exclude cardiomyocyte lipogenesis, as previously described.39

In the present work, we investigated whether the non-contractile cardiac stromal compartment is the cellular source contributing to excess adipocytes in ACM hearts. Cardiac mesenchymal stromal cells are the most abundant cell type in the heart, have a mesenchymal origin, and exert a fundamental role in maintaining cardiac structural and functional homeostasis in physiologic and pathologic conditions.29 Importantly, upon appropriate culture conditions C-MSC can differentiate into adipocytes.27 Therefore we hypothesized and demonstrated that when affected by ACM mutations, C-MSC enhance their ability to accumulate lipids, differentiate into adipocytes, and play a role in fibro-fatty substitution.

Based on the obtained evidence, we advocate C-MSC as a novel non-contractile cell type contributing to excess adipocyte accumulation in ACM. PKP2-silenced murine epicardial cells were so far the best characterized non-contractile cell type as a putative source of lipid accumulation.35,40 Epicardial cells have been recognized as second heart field-derived progenitors of a large fraction of non-myocyte heart cells, including fibroblasts.41 Epicardial cells silenced for PKP2 were shown to be able to accumulate more lipid than control epicardial cells (EPC).35 We believe that this observation is not conflicting with our results. Epicardial cells may undergo epithelial-mesenchymal transition, thus retaining, as adult MSC, the ability to accumulate lipid in a PKP2-dependent manner.

Resident cardiac stem cells (c-KIT/SCA1) from PG transgenic mice have been indicated by Lombardi et al.20 as a source of adipocytes.

**Figure 5** Cardiac mesenchymal stromal cells obtained from arrhythmogenic cardiomyopathy patients (ACM), cultured in adipogenic medium, accumulate more lipid droplets than control cardiac mesenchymal stromal cells (NON-ACM). Representative images of cardiac mesenchymal stromal cells from bioptic samples of patients affected and not affected by arrhythmogenic cardiomyopathy, cultured for 72 h and 1 week in adipogenic medium, stained with Oil Red O (ORO; n = 6 vs. n = 4). The scale bar indicates 50 μm. Quantification of the luminance of the 255 red staining is reported on the right: intensity is expressed in arbitrary units.
We also found occasional c-KIT positive cells undergoing adipogenic differentiation in ACM patients' hearts; however, in our opinion, the rates were too low to account for a major role in lipid accumulation. We have previously shown that c-KIT positive cells can express mesenchymal markers, but they represent only the 0.5–1.5% of isolated C-MSC (see Supplementary material online, Figure S1 and Table S2). Therefore, the diffuse, homogeneous adipogenic differentiation of the whole ACM C-MSC population observed in vitro argues against a major direct contribution of cardiac progenitor cells in the development of fibro-adipose substitution in ACM.

Plakoglobin predominant nuclear staining in cultured C-MSC is consistent with previous findings in other cell types, even if this phenomenon is still controversial in vivo. Therefore, the diffuse, homogeneous adipogenic differentiation of the whole ACM C-MSC population observed in vitro argues against a major direct contribution of cardiac progenitor cells in the development of fibro-adipose substitution in ACM.

Figure 6 Arrhythmogenic cardiomyopathy cardiac mesenchymal stromal cells in adipogenic medium show higher transcription of adipogenic genes and proteins and a lower transcription of anti-adipogenic and proliferation genes, compared to control cells. ACM: samples from patients affected by arrhythmogenic cardiomyopathy; NON-ACM: samples from patients not affected by arrhythmogenic cardiomyopathy. (A) Comparison of transcript abundance (genes threshold cycles [Ct] with respect to the house-keeping gene GAPDH) between RNA extract of cardiac mesenchymal stromal cells from patients not affected and affected by arrhythmogenic cardiomyopathy (n ≥ 4), after 72 h culture in adipogenic medium. (B) ADIPOQ and PLIN1 gene expression positively correlates with lipid accumulation (Oil Red O staining) in arrhythmogenic cardiomyopathy samples. (C) Comparison of PLIN1, PPARγ, and FABP4 protein abundance between protein extracts of cardiac mesenchymal stromal cells from patients not affected by arrhythmogenic cardiomyopathy (n = 3) and from arrhythmogenic cardiomyopathy patients (n = 5), after 72 h culture in adipogenic medium. Quantification of the mean protein abundance relative to GAPDH, and normalized for values of samples from patients not affected by arrhythmogenic cardiomyopathy, is shown in the graphs.

We also found occasional c-KIT positive cells undergoing adipogenic differentiation in ACM patients' hearts; however, in our opinion, the rates were too low to account for a major role in lipid accumulation. We have previously shown that c-KIT positive cells can express mesenchymal markers, but they represent only the 0.5–1.5% of isolated C-MSC (see Supplementary material online, Figure S1 and Table S2). Therefore, the diffuse, homogeneous adipogenic differentiation of the whole ACM C-MSC population observed in vitro argues against a major direct contribution of cardiac progenitor cells in the development of fibro-adipose substitution in ACM.

Plakoglobin predominant nuclear staining in cultured C-MSC is consistent with previous findings in other cell types, even if this phenomenon is still controversial in vivo. The partial rescue of the lipid accumulation phenotype in C-MSC treated with BIO is suggestive of WNT pathway involvement in ACM, although not to the exclusion of the contributions from other mechanisms.

Notably, C-MSC may be viewed as a novel cell platform for studying ACM in vitro. Different cellular models of disease pathogenesis have been previously proposed, mainly focused on the contractile component of the heart: PKP2-silenced cardiomyocytes, murine DSP-deficient HL1 or mouse cardiomyocytes, PG transgenic mouse c-KIT/SCA1 cells, and recently different lines of iPSC-derived cardiomyocytes. However, these models have specific intrinsic problems, including the lack of endogenous genetic mutation, the non-human cell origin, possible reprogramming biases, or unsuitability for high-throughput screenings. Mutated and control C-MSC are primary cells directly obtained from human ventricular tissue, easy to isolate and amplify, carrying the whole patient's genetic setting (ACM mutation and background). Therefore, we think that C-MSC are suitable tools for disease modelling.

Notwithstanding the relevance of substrate insults in generating or worsening the arrhythmic phenotype of ACM, recent reports highlight ACM as a primary electrical cardiac disease. Although we have observed that in growth conditions Cx43 expression levels and cellular localization (see Supplementary material online, Figure S8) are similar in ACM and NON-ACM C-MSC, future studies are needed to establish whether differentiating C-MSC can contribute to modulate cell survival/death, as well as functional properties of cardiomyocytes, and of other cardiac cell types.

In conclusion, the present article reports unprecedented evidence that C-MSC are a source of adipocytes in ACM heart tissue and that ACM-derived primary C-MSC differentiate into adipocytes in vitro.
Study limitations

The variability we observed among cells derived from different patients, both in lipid accumulation and in gene expression, may at least partially correlate with patient clinical presentation and genetic status. Though limited by the small sample size, an analysis of C-MSC ORO staining intensities stratified by genotype indicated that cells from PKP2 mutation carriers have a higher predisposition to lipid accumulation than C-MSC from patients without mutations in known ACM-related genes ($P = 0.058$). Further, all PKP2 mutation carriers biopsies reached the major diagnostic criterion of substantial fibro-fatty replacement of myocardium. These observations, if validated in studies with larger cohorts, may complete arrhythmic phenotype–genotype correlations reported by Bao and colleagues, who demonstrated that mutation carriers (in particular of PKP2) are subject to spontaneous and induced VT more often than patients without mutations.

We obtained C-MSC only from PKP2 mutation carriers or from patients with no mutations in known ACM-associated genes (Table 1). Therefore, our results need to be tested in patients with other desmosomal or non-desmosomal mutations.
Cardiac mesenchymal stromal cells cultured in adipogenic medium show PLIN1 staining similar to differentiating cells in patients’ hearts. Not having access to LV bioptic samples of ACM patients, we were unable to explore possible differences in adipogenic propensity between the right and left ventricles.

Supplementary material
Supplementary material is available at European Heart Journal online.

Author contributions
E.S. and G.I.C.: performed statistical analysis; G.P. and C.T.: handled funding and supervision; E.S., S.B., C.C., E.G., V.M., A.D.R., F.M.F., V.C., G.P., M.C., I.S., E.C., N.O.A., and C.P.: acquired the data; E.S., G.P., M.C.C., and A.R.: conceived and designed the research; E.S., S.B., and C.C.: drafted the manuscript; M.C., A.P., G.d’A., G.I.C., A.R., M.C.C., and A.R.: made critical revision of the manuscript for key intellectual content.

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References
1. Asimaki A, Kleberger AG, Saffitz JE. Pathogenesis of arrhythmogenic cardiomyopathy. Con J Cardiol 2015. doi: 10.1016/j.cjca.2015.04.012.
2. Cerrone M, Noorman M, Lin X, Chiorozi H, Liang FX, van der Nagel R, Hund T, Birchmeier W, Mohler P, van Veen TA, van Rijen HV, Delmar M. Sodium current deficit and arrhythmogenesis in a murine model of plakophilin-2 haploinsufficiency. Cardiovasc Res 2012;95:460–468.
3. Sato PY, Musa H, Coombs W, Guerrero-Serna G, Patino GA, Taffet SM, borm LS, Delmar M. Loss of plakophilin-2 expression leads to decreased sodium current and slower conduction velocity in cultured cardiac myocytes. Circ Res 2009;105:523–526.
4. Oxford NM, Evertt M, Coombs W, Fox KR, Kraus M, Gelzer AR, Saffitz J, Taffet SM, Moise NS, Delmar M. Molecular composition of the intercalated disc in a spontaneous canine animal model of arrhythmogenic right ventricular dysplasia/cardio-myopathy. Heart Rhythm 2007;4:1196–1205.
5. Swope D, Cheng L, Gao E, Li J, Radice GL. Loss of cadherin-binding proteins beta-catenin and plakoglobin in the heart leads to gap junction remodeling and arrhythmogenesis. Mol Cell Biol 2012;32:1056–1067.
6. Gomes J, Finlay M, Ahmed AK, Caccio EF, Asimaki A, Saffitz JE, Quarta G, Nobles M, Sypiris P, Chaubey S, McKenna WJ, Tinker A, Lambaze PD. Electrophysiological abnormalities precede overt structural changes in arrhythmogenic right ventricular cardiomyopathy due to mutations in desmoplakin II combined murine and human study. Eur Heart J 2012;33:1942–1953.
7. Norgett EE, Hatzell SS, Carvalho-Huerta L, Cabezas JC, Common J, Purks PE, Whitstone N, Leigh IM, Stevens HF, Kessell DP. Recessive mutation in desmoplakin disrupts desmoplakin-intermediate filament interactions and causes dilated cardiomyopathy, woolly hair and keratoderma. Hum Mol Genet 2000;9:2761–2766.
8. McKoy G, Protonotarios N, Crosby A, Tsatsopoulou A, Anastassakis A, Cooran A, Norman M, Baboonian C, Jeffrey S, McKenna WJ. Identification of a deletion in plakoglobin in arrhythmogenic right ventricular cardiomyopathy with palmoplantar keratoderma and woolly hair (Naxos disease). Lancet 2000;355:2119–2124.
9. Gerull B, Heuser A, Wichter T, Paul M, Basson CT, McDermott DA, Lerman BB, Markowitz SM, Elinor PT, MacRae CA, Peters S, Grossmann KS, Drenckhahn D, Michely B, Sasse-Klaassen S, Birchmeier W, Dietz R, Breithardt G, Schulze-Bahr E, Thierfelder L. Mutations in the desmosomal protein plakoglobin are common in arrhythmogenic right ventricular cardiomyopathy. Nat Genet 2004;36:1162–1164.
10. Pilichou K, Nava A, Basso C, Beffagna G, Bause B, Lorenzon A, Frigo G, Vettori A, Vallette M, Towlbin J, Thieme G, Danielli GA, Rampazzo A. Mutations in desmoglein-2 gene are associated with arrhythmogenic right ventricular cardiomyopathy. Circulation 2006;113:1171–1179.

Figure 8: Cardiac mesenchymal stromal cells cultured in adipogenic medium show PLIN1 staining similar to differentiating cells in patients’ hearts. (A) Immunostaining for PLIN1 of cells from patients not affected by arrhythmogenic cardiomyopathy (NON-ACM) and from arrhythmogenic cardiomyopathy patients (ACM), cultured in adipogenic medium for 72 h. The scale bar indicates 10 μm. (B) Immunostaining for PLIN1 on FFPE sections of an arrhythmogenic cardiomyopathy (ACM) explanted heart. The scale bar indicates 20 μm. Nuclei are stained with Hoechst 33258.
11. Syrris P, Ward D, Evans A, Asnaki A, Gandjbakhch E, Sen-Chowdhry S, McKenna WJ. Arrhythmogenic right ventricular dysplasia/cardiomyopathy asso-
ciated with mutations in the desmosomal gene desmocollin-2. Am J Hum Genet
2006;79:978–984.
12. Johnson JL, Najor NA, Green KJ. Desmosomes: regulators of cellular signaling
and adhesion in epidermal health and disease. Cold Spring Harb Perspect Med 2014;
4 a015297.
13. Lazzarini E, Jongbloed JD, Piliouch K, Thiene G, Basso C, Bikhler K, Charbon B,
Swertz M, van Tintelen JP, van der Zwaag PA. The ARVDIC genetic variants data-
base: 2014 update. Hum Mutat 2015;36:403–410.
14. Chen X, Bonne S, Hatzfeld M, van Roy F, Green KJ. Protein binding and functional
characterization of plakophilin 2. Evidence for its diverse roles in desmosomes and
beta -catenin signaling. J Biol Chem 2002;277:10512–10522.
15. Cheng W, Li B, Kajstura J, Li P, Wolin MS, Sonnenblick EH, Hintze TH, Olivetti G,
Anversa P. Stress-stretched-induced myocyte cell death. J Clin Invest 1995;96:
2247–2259.
16. Li J, Swope D, Raes N, Cheng L, Muller EJ, Radice GL. Cardiac tissue-restricted
deletion of plakoglobin results in progressive cardiomyopathy and activation of
(beta -catenin signaling. Mol Cell Biol 2011;31:1134–1144.
17. Chen SN, Gurha P, Lombardi R, Ruggiero A, Willerson JT, Marian AJ. The hippo
pathway is activated and is a causal mechanism for adipogenesis in arrhythmogenic
cardiomyopathy. Circ Res 2014;114:454–468.
18. Ross SE, Hemati N, Longo KA, Bennett CN, Lucas PC, Erickson RL, MacDougald
OA. Inhibition of adipogenesis by Wnt signaling. Science 2000;289:
950–953.
19. Garcia-Gras E, Lombardi R, Giordano C, Gallo P. Myocyte transdifferentiation:
a possible pathogenic mechanism for arrhythmogenic right ventricular cardiomyop-
yopathy. J Clin Invest 2006;116:2012–2021.
20. Lombardi R, da Graca-Cabreira-Hansen M, Basso A, Fromm RR, Willerson JT,
Marian AJ. Nuclear plakoglobin is essential for differentiation of cardiac progenitor
cells to adipocytes in arrhythmogenic right ventricular cardiomyopathy. Circ Res
2013;110:1342–1353.
21. d’Amati G, di Gioia CR, Giordano C, Gallo P. Possible pathogenic mechanism for
arrhythmogenic right ventricular cardiomyopathy. Arch Pathol Lab Med 2000;124:
287–290.
22. Ma D, Wei H, Lu J, Hao S, Zhang G, Sun X, Oh Y, Tan SH, Lu J, Song L, He X.
Correlation of ventricular arrhythmias with genotype in arrhythmogenic
right ventricular cardiomyopathy. J Obes (Lond) 2015;2015:1076–1084.
23. Caspi O, Huber I, Gepstein A, Arbel G, Mazels L, Boulou M, Gepstein L. Modeling
of arrhythmogenic right ventricular cardiomyopathy with human induced pluripo-
tent stem cells. Circ Cardiovasc Genet 2013;6:557–568.
24. Lombardi R, Dong J, Rodriguez G, Basso A, Leung TK, Schwartz RJ, Willerson JT,
Brugada R, Marian AJ. Genetic fate mapping identifies second heart field progenitor
cells as a source of adipocytes in arrhythmogenic right ventricular cardiomyopathy.
Circ Res 2009;104:1076–1084.
25. Piliouch K, Bezizza CR, Thiene G, Basso C. Arrhythmogenic cardiomyopathy: trans-
genic animal models provide novel insights into disease pathology. Circ Cardi-
ovas Genet 2011;4:318–326.
26. da Silva Meirelles L, Chagastelles PC, Nardi NB. Mesenchymal stem cells reside in
virtually all post-natal organs and tissues. J Cell Sci 2006;119:2204–2213.
27. Rossini A, Frati C, Lagrassa G, Graiani G, Scapece A, Cavalli S, Musso E, Baccarini M,
Di Segni M, Fagnoni F, Germani A, Quaini E, Marz Y, Xu Q, Barbuti A, DiFrancesco D,
Pompilio G, Quaini F, Gaetaano C, Capogrossi MC. Human cardiac and bone marrow
stromal cells exhibit distinctive properties related to their origin. Cardiovasc Res 2010;89:
650–660.
28. Jürgutis B. Ventricular remodeling after infarction and the extracellular collagen
matrix: when is enough enough? Circulation 2003;108:1395–1403.
29. Brown RD, Amblor SK, Mitchell MD, Leng CS. The cardiac fibroblast: therapeutic
target in myocardial remodeling and failure. Annu Rev Pharmacol Toxicol 2005;45:
657–687.
30. Pino AM, Rosen CJ, Rodriguez J. In osteoporosis, differential expression of mesenchymal
stem cells (MSCs) improves bone marrow adipogenesis. Biol Res 2012;45:
279–287.
31. Gambini E, Pesce M, Persico L, Bassetti B, Gambini A, Alamanni F, Agrifoglio M,
Capogrossi MC, Pompilio G. Patient profile modulates cardiac c-kit (+) progenitor
cell availability and amplification potential. Transl Res 2012;160:363–373.
32. Sun Z, Gong J, Wu X, Wu W, Lu X, Diao J, Wu JW, Yang H, Yang M, Li P. PeriPP1
promotes unilocular lipid droplet formation through the activation of
Pp27 in adipocytes. Nat Commun 2013:41594.
33. Marcus FI, McKenna WJ, Sherrill D, Basso C, Bache B, Blumenke DA, Calcins H,
Corrado D, Cox MG, Daubert JP, Fontaine G, Gear K, Hauer R, Nava A, Picard MH,
Protonatorios N, Saffitz JE, Sanborn DM, Steinberg JS, Tandri H, Thieme G, Töwinja
TA, Tsatsopoulos A, Wichter T, Zareba W. Diagnosis of arrhythmogenic
right ventricular cardiomyopathy/dysplasia: proposed modification of the
Task Force Criteria. Eur Heart J 2010;31:806–814.
34. Luo Q, Kang Q, Si W, Jiang W, Park JK, Peng Y, Li X, Liu H-H, Luo J, Montag AG,
Haydon RC, He TC. Connective tissue growth factor (CTGF) is regulated by
Wnt and bone morphogenetic proteins signaling in osteoblast differentiation of
mesenchymal stem cells. J Biol Chem 2004;279:55958–55968.
35. Matthews SA, Tafset S, Delmar M. Plakophilin-2 and the migration, differentiation
and transformation of cells derived from the epicardium of neonatal rat hearts. Cell
Commun Adhes 2011;18:73–84.
36. Delva E, Tucker DK, Kowalczyk AP. The desmosome. Cold Spring Harb Perspect Biol
2009;1:a002543.
37. Kim C, Wong J, Wen J, Wang S, Wang C, Spiering S, Kang NG, Suri PL, Leone TC,
Marine JE, Calcins H, Kelly DP, Judge DP, Chen HS. Studying arrhythmogenic
right ventricular dysplasia with patient-specific iPSCs. Nature 2013;494:
105–110.
38. Robertson C, Tran DD, George SC. Concise review: maturation phases of human
pluripotent stem cell-derived cardiomyocytes. Stem Cells Dev 2013;21:
829–837.
39. Djouadi F, Lecarpentier Y, Hebert JL, Charron P, Bastin J, Caillet A. A potential
link between peroxisome proliferator-activated receptor signaling and the patho-
genesis of arrhythmogenic right ventricular cardiomyopathy. Cardiovasc Res 2009;
84:83–90.
40. Oxford EM, Musa H, Maass K, Coombs W, Taffet SM, Delmar M. Connexin43 re-
modeling caused by inhibition of plakophilin-2 expression in cardiac cells. Circ Res
2007;101:703–711.
41. Fishman MC, Chien KR. Fashioning the vertebrate heart: earliest embryonic deci-
sions. Development 1997;124:2099–2117.
42. Gambini E, Pompilio G, Biondi A, Alamanni F, Capogrossi MC, Agrifoglio M,
Pesce M. C-kit + cardiac progenitors exhibit mesenchymal markers and preferen-
tial cardiovascular commitment. Cardiovasc Res 2010;89:362–373.
43. Zhang Z, Stroud MJ, Zhang J, Fang X, Ouyang K, Kimura K, Mu Y, Dalton ND, Gu Y,
Bradford WH, Peterson KL, Cheng H, Zhou X, Chen J. Normalization of Naxos
plakoglobin levels restores cardiac function in mice. J Clin Invest 2015;125:
1708–1712.
44. Noorman M, Hakim S, Kessler E, Groeneweg J, Cox MG, Asimaki A, van Rijen HV,
Noorman M, Hakim S, Kessler E, Groeneweg J, Cox MG, Asimaki A, van Rijen HV,
van Stuijvenberg L, Chilouris H, van der Heyden MA, Vos MA, de Jonge N, van
demmaizer, W. Ann Surg, 1998;197:1724–1728.
45. Gambini E, Pompilio G, Quaini F, Gaetaano C, Capogrossi MC. Human cardiac
and bone marrow stromal cells exhibit distinctive properties related to their origin.
Cardiovasc Res 2010;89:650–660.
46. Bao J, Wang J, Yao Y, Wang Y, Fan X, Sun K, He DS, Marcus H, Zhang S, Hui R,
Song L. Correlation of ventricular arrhythmias with genotype in arrhythmogenic
right ventricular cardiomyopathy. Circ Cardiovasc Genet 2013;6:552–556.