Human iPSC-Derived Cardiomyocytes of Peripartum Patients With Cardiomyopathy Reveal Aberrant Regulation of Lipid Metabolism

Peripartum cardiomyopathy (PPCM) is a rare form of heart failure occurring in the last trimester of pregnancy or in the first months after delivery.¹ The diagnosis is based on exclusion criteria, and specific biomarkers remain unidentified.¹ Proposed mechanisms of PPCM pathophysiology are based on various mouse models harboring cardiac-specific genetic defects,²,³ which demonstrated typical PPCM characteristics, but mutations in these genes remain to be associated with PPCM in patients.⁴ This study aims to identify aberrant pathways in cardiomyocytes obtained from induced pluripotent stem cells (iPSC) derived from patients with PPCM.

Two patients with PPCM with acute heart failure (diagnosed within weeks after delivery) have been included in the study. Patients A and B were aged 37 and 28 years with a left ventricular ejection fraction of 17% and 45% and elevated levels of N-terminal pro-B-type natriuretic peptide 1309 ng/L and 2743 ng/L at presentation, respectively. Patient A was compared with her sister and patient B with her mother in paired analyses; both controls were healthy, had a normal echocardiogram, and have had multiple uncomplicated deliveries. Patients were screened for genetic variations in 61 genes associated with cardiomyopathies, but none was identified. All participants provided written informed consent. This study was approved by the local Medical Ethical Committee (METc:2014.104).

Pregnancy-associated wall stress was simulated in vitro by applying cyclic mechanical stretch to the iPSC-derived cardiomyocytes (iPSC-CM; Figure [A]). RNA sequencing analysis of these conditions resulted in a total of 3316 differentially expressed genes (Figure [B]). We identified 95 common differentially expressed genes in healthy mechanically stretched iPSC-CM and in both static and stretched PPCM iPSC-CM compared with healthy static iPSC-CM (designated as the intersection of interest in yellow, Figure [B]). Therefore, a stretch-induced stress response was observed in all mechanically stretched cells, but also in static PPCM iPSC-CM. Relative expression levels of these 95 genes were visualized in a heat map (Figure [C]), and a pathway overrepresentation analysis was performed. Predicted gene ontology terms were organized into superclusters (Figure [D]). Gene ontology terms pertained mostly to lipid metabolism as part of the fatty alcohol biosynthesis supercluster of gene ontology terms, and 28 of 95 differentially expressed genes were associated to metabolic processes (depicted in red, Figure [C]). In silico transcription factor enrichment analysis identified key transcription factors SREBP1 (sterol regulatory element-binding transcription factor 1), NFY (nuclear transcription factor Y), SP1 (Sp1 transcription factor), MAX (MYC-associated factor X), and CEBPB (CCAAT enhancer-binding protein beta).

To underscore the relevance of these findings, we aimed to validate our findings in an established PPCM mouse model caused by cardiac-specific STAT3 conditional knockout (STAT3-CKO).² These STAT3-CKO mice consistently developed PPCM as demonstrated by a severely reduced cardiac function (Figure [E]). Gene expression analysis

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**Figure.** Pathways related to lipid metabolism were aberrantly regulated in PPCM-specific iPSC-derived cardiomyocytes.

A. Schematic overview of the experimental setup. Multiple iPSC clones were generated from patient A (4 clones), patient B (2 clones), and both controls (2 clones from each). Each clone has been differentiated to cardiomyocytes at least 3 times. Cardiomyocytes have been subjected to cyclic equiaxial mechanical stretch (15% elongation at 1 Hz for 48 hours) to mimic pregnancy-related hemodynamic stress.

B. Venn diagram denoting the number of differentially expressed genes (DEGs) in stretched healthy cardiomyocytes, static and stretched cardiomyocytes derived from patients with PPCM, and reciprocal intersections compared with static healthy cardiomyocytes (background). The yellow intersection of interest marks the overlapping set of 95 DEGs in cardiomyocytes (Continued).
of left ventricular tissue of postpartum STAT3-CKO mice (ie, after 2 pregnancies and nursing periods) showed that key metabolic genes were also differentially regulated compared with nulliparous STAT3-CKO mice (Figure [F]).

Analysis of metabolic substrate utilization in the PPCM iPSC-CM and isolated STAT3-CKO cardiomyocytes was performed by means of a Seahorse assay–based Mito Fuel Flex test. Cytosolic anaerobic glycolysis was markedly blunted in PPCM iPSC-CM at baseline and after inhibition of lipid metabolism (Figure [G]). To assess substrate dependence, we measured cellular viability after specific inhibition of metabolic pathways (Figure [H]). Viability was rapidly decreased by inhibition of total lipid metabolism, total glucose metabolism, or both within the first hour. Interestingly, healthy iPSC-CM recovered over time, which was not observed in PPCM iPSC-CM after 48 hours. Inhibition of glucose metabolism resulted in elevated OCR in isolated wild-type cardiomyocytes, which was blunted in STAT3-CKO cardiomyocytes; this did not affect cytosolic anaerobic glycolysis (Figure [I]).

Under physiological circumstances, maternal lipid metabolism is increased during the last trimester of pregnancy and quickly normalizes after delivery.5 This metabolic transition is greatly dependent on transcription factors governing lipid metabolism. Our results demonstrate that iPSC-CM derived from patients with PPCM showed disrupted regulation of pathways related to lipid metabolism, which was validated in an established PPCM mouse model. Furthermore, we observed impaired anaerobic glycolysis in PPCM iPSC-CM after inhibition of lipid metabolism. Healthy iPSC-CM demonstrated adequate metabolic plasticity by activation of glucose utilization in response to inhibition of lipid metabolism; this metabolic switch was blunted in PPCM iPSC-CM. Isolated wild-type cardiomyocytes immediately switched to lipid metabolism after inhibition of glucose metabolism, which was not observed in STAT3-CKO cardiomyocytes.

Although the number of patients included in this study remains a limitation, this study highlights that commonly affected pathways could be identified in 2 different families. STAT3 and PGC1α were not identified as differentially expressed genes in any condition, but these might be altered posttranscriptionally in patients with PPCM and may exacerbate PPCM progression.5 However, patients with PPCM may have mutations or epigenetic alterations in other genes, which may explain altered cardiac metabolism in PPCM iPSC-CM. Moreover, the correlation between parity and PPCM severity remains to be further investigated.

In conclusion, our data show that lipid metabolism is widely affected in PPCM iPSC-CM and highlight the potential role of metabolic regulation as a key factor for PPCM susceptibility.

ARTICLE INFORMATION

Data sharing: All data and materials have been made publicly available at Array Express and can be accessed at https://www.ebi.ac.uk/arrayexpress/experiments/E-MTAB-9053/. All cell lines used in this study have been registered in the Human Pluripotent Stem Cell Registry at https://hpscreg.eu/.

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Disclosures
None.

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