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Chapter 7:
Serum supplemented culture medium masks hypertrophic phenotypes in human pluripotent stem cell derived cardiomyocytes

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Abstract
It has been known for over 20 years that fetal calf serum can induce hypertrophy in cultured cardiomyocytes but this is rarely considered when examining cardiomyocytes derived from pluripotent stem cells (PSC). Here, we determined how serum affected cardiomyocytes from human embryonic- (hESC) and induced pluripotent stem cells (hiPSC) and hiPSC from patients with hypertrophic cardiomyopathy linked to a mutation in the MYBPC3 gene. We first confirmed previously published hypertrophic effects of serum on cultured neonatal rat cardiomyocytes evidenced as increased cell surface area and beating frequency. We then found that serum increased the cell surface area of hESC- and hiPSC-derived cardiomyocytes and their spontaneous contraction rate. Phenylephrine, which normally induces cardiac hypertrophy, had no additional effects under serum conditions. Likewise, hiPSC-derived cardiomyocytes from 3 MYBPC3 patients which had a greater surface area than controls in the absence of serum as predicted by their genotype, did not show this difference in the presence of serum. Serum can thus alter the phenotype of human PSC derived cardiomyocytes under otherwise defined conditions such that the effects of hypertrophic drugs and gene mutations are underestimated. It is therefore pertinent to examine cardiac phenotypes in culture media without or in low concentrations of serum.
Introduction

Human pluripotent stem cells (hPSC), particularly those derived as induced pluripotent stem cells (hiPSC) from patients with genetic diseases, are increasingly regarded as useful for disease modeling and drug target discovery. hiPSC are considered particularly valuable because they capture the genome of the individual from whom they are derived. They have the ability to self-renew over long periods and differentiate into all cell types of the body and thus represent a permanent resource for modeling human disease. Although directed differentiation is still challenging for many lineages, cardiomyocytes were among the first functional cells to be derived and characterized from human embryonic stem cells (hESC) [1-3] and they can now be generated efficiently from hiPSC using similar protocols. Cardiomyocytes derived from human (h)PSC have been successful in mimicking reported drug responses in patients [4-7] including the positive chronotropic response to phenylephrine (PE), an α-adrenergic agonist, and isoproterenol (ISO), a β-adrenergic agonist [3]. These adrenergic stimuli have also been shown to induce pathological hypertrophy in cultured cardiomyocytes [8-10]. Pathological hypertrophy is an abnormal increase in cell size accompanied by the re-expression of fetal cardiac genes [11]. Aside from drug-induced cardiac hypertrophy, cardiomyocytes from hiPSC of patients with mutations associated with hypertrophic cardiomyopathy (HCM) also show features of hypertrophy and are thus potentially useful disease models [12]. Cardiomyocytes from patient hiPSC have already increased our knowledge of cardiac diseases [12-14], complementing and extending previous work based on clinical data and primary human cardiomyocyte cultures.

Whilst the uses of hPSC-derived cardiomyocytes are becoming well established, there is still no single method to derive these cells in culture. Although differentiation protocols are increasingly based on defined media and timed growth factor addition [15-17; reviewed in 18], fetal calf serum (FCS) is still often present, either during differentiation or during long-term maintenance to support survival as the cardiomyocytes attain some degree of maturity. Of 119 articles published using hPSC-CM in 2012 and 2013, 54 used 5% or higher concentrations of serum in the differentiation and/or maintenance medium. In others serum was present but at concentrations ≤2%. Since the exact composition of serum is unknown and varies considerably from batch to batch [19], its presence can confound interpretation of experiments investigating disease phenotype or drug responses. For example, it has long been known that serum or serum deprivation can profoundly affect primary neonatal rat cardiomyocytes (Rat-CM) in culture [20-24]. Cell size and protein-to-cell ratio can increase and additional stress-fiber-like structures may be induced in response to serum, which is indicative of cardiac hypertrophy [21, 24]. Other studies have used serum as a hypertrophic stimulus for cardiomyocytes [20, 25, 26], in place of the more commonly used hypertrophic stimuli PE or ISO [10]. The impact of serum on the cardiomyocyte phenotype and, more specifically, on the manifestation of the disease in the case of cardiomyocytes derived from patient hiPSC, remains unreported.

Here we examined the effects of serum and PE in the widely used Rat-CM model and confirmed findings reported previously. We then investigated the effects of serum on healthy- and HCM patient-derived hiPSC-CM obtained under serum-free, defined conditions. Parameters examined included cell surface area, sarcomeric structure and beating frequency. These are common descriptors of phenotype in the literature. The effects of serum were compared with those induced by the hypertrophic drugs PE and ISO. We found that serum induced increased cell surface area and beating frequency and disrupted sarcomeric structure, indicating a hypertrophic response in both hESC-CM and hiPSC-CM. This masked the disease-associated hypertrophy of HCM-hiPSC-CM that was evident in the absence of serum. These results have important implications for conditions under which disease phenotypes are examined.
Chapter 7

Material and Methods

Ethics statement

The isolation of neonatal rat cardiomyocytes was approved by the Animal Experiment Committee of Leiden University Medical Center and complies with the Guide for the Care and Use of Laboratory Animals as stated by the US National Institutes of Health. Human skin biopsies were obtained from patients after individual written permission using standard informed consent procedures following approval for use in this study by Leiden University Medical Center’s medical ethics committee; this conforms to the Declaration of Helsinki. Control skin samples were obtained as waste tissue from donors in accordance with the Dutch federation of Biomedical Scientific Societies “Use of human tissue for scientific research” and “Code of good use” directives. All samples were collected by the treating physician and then anonymized.

Neonatal rat cardiomyocytes

Neonatal rat cardiomyocytes were isolated as previously described [27]. Briefly, rats were anesthetized with inhaled isoflurane (4-5%) and adequate anesthesia was confirmed by the absence of pain reflexes before excising the hearts. Ventricles from the dissected hearts were minced and dissociated using collagenase and DNase and then suspended in Ham’s F10 medium (ICN Biomedicals) with 10% horse serum (Invitrogen) and 10% FCS. To allow for preferential attachment and negative selection of the non-cardiomyocytes, the cell suspension was pre-plated for 1h in a tissue culture dish. The non-adherent cardiomyocytes were then collected and plated on Matrigel coated coverslips (plating density of 20,000 cells/cm2) in isolation medium or serum-free BEL medium (Iscove’s Modified Dulbecco’s Medium (IMDM) supplemented with L-glutamine and 25mM HEPES (Invitrogen), F12 Nutrient Mixture (HAM) supplemented with Glutamax (Invitrogen), 5% protein free hybridoma medium (Invitrogen), 0.25% deionized albumin from BSA (Sigma,) in IMDM, 1% Chemically Defined Lipid Concentrate (Invitrogen),

Figure 1) Responses of neonatal Rat-CM to serum. A) Schematic timeline of treatment procedure. Percentages indicate the serum concentrations. In parallel controls, PE was omitted. B) Cell surface area of Rat-CM in 20%, or 0% serum at 12h, 36h and 60h after plating and 5% serum 36h and 60h after plating. *=p<0.05 20% serum (12h) vs. other conditions (n≥67). C) Cell surface area of neonatal Rat-CM (60h) maintained in isolation medium (20% serum) throughout or subsequently reduced to 5% or 0% serum, or initially plated in 0% serum ±100 μM PE for 24 h. *=p<0.05 untreated vs. PE-treated (n≥59 cells). D) Beating frequency of Rat-CM (60h) ± 100 μM PE. *=p <0.05 untreated vs. PE-treated (n≥7 measurements).
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0.1% Insulin-Transferrin-Selenium-X supplement (Invitrogen), 450 μM 1-Thioglycerol (Sigma), 5mg/ml L-ascorbic acid 2-phosphate (Sigma), 1% Glutamax (Invitrogen), 25 U/ml penicillin, 25 μg/ml streptomycin (both Invitrogen). 12 h later the medium was replaced and the cells were maintained in the original isolation medium (20% serum), 5% FCS in BEL or serum-free medium BEL for 24 h before treatment with 100 μM PE for an additional 24 h.

hESC culture

The Nkx 2-5eGFP/w hESC line [28] was maintained as a single cell culture in Dulbecco’s Modified Eagle Medium (DMEM)/F12 (Invitrogen) containing 20% Knockout Serum replacement (Invitrogen), 1x MEM-Non-Essential amino acids (Invitrogen) 0.1 mM 2-Mercaptoethanol (Invitrogen), 10 ng/ml Human basic fibroblast growth factor (bFGF) (Peprotech) , 25 U/ml penicillin and 25 μg/ml streptomycin (both Invitrogen) and passaged every 3-4 days, as previously described [29].

Human iPSC Derivation

Human dermal fibroblasts were isolated from biopsies from a control subject and patients carrying a MYBPC3 gene mutation and displaying an HCM phenotype. Fibroblasts were reprogrammed to hiPSC as previously described [30]. hiPSC were routinely cultured on Matrigel (BD Biosciences)-coated tissue culture dishes in mTESR according to the manufacturer’s protocol (Stem Cell Technologies). The cells were mechanically passaged weekly using 1 mg/ml Dispase (Gibco).

Transgene removal and confirmation

Based on morphology, hESC-like colonies were manually picked 30 days after transduction. Cells were adapted to enzymatic culture on mouse embryonic feeder cells [29] and then transfected with pLV.hCMV-IE.FLP.e. PurR.hHBVPRE [31] to remove the reprogramming cassette. Removal was confirmed by PCR of the FRT region (triple primer set): 5’-CGAGTCGGATCTCCCTTTGGGC-3’, 5’TGGAAAGGGCTACGTAGCTAGC-3’, 5’-GGTTCCCTAGTTAGCCAGAGAGC-3’. Transgene-free clones were readapted to mechanical passage on Matrigel with mTESR. Frozen stocks were made at passage 5 and cells were used for experimentation between passage 10 and 30. LUMC0004iCtrl (Healthy), LUMC0033iMyBPC (HCM1), LUMC0034iMyBPC (HCM2), LUMC0035iMyBPC (HCM3) hiPSC were used in this study.

Generation of Cardiomyocytes

Cardiomyocytes were generated from the Nkx 2-5eGFP/w hESC line [28] and hiPSC using a monolayer method in defined medium with timed addition of growth factors and small molecules. hESC were plated at a density of 10,000 cells/cm² and hiPSC were dissociated into small clumps three days prior to differentiation and allowed to attach on Matrigel coated dishes in mTESR (~1.5 clumps/cm²) before starting the differentiation procedure. Differentiation was induced in low-insulin (1mg/L), “BEL medium” [16] plus 20 ng/ml Activin A (R&D systems), 20 ng/ml BMP4 (R&D systems) and 1.5 μM CHIR99021 (Axon). These factors were removed after 3 days and replaced with BEL plus 5 μM XAV939 (R&D systems) and Matrigel (BD bioscience) (1:100) to prevent cell detachment. The cells were refreshed with BEL three days later and subsequently maintained in BEL (changing medium twice a week). On average the differentiation efficiency of this method is 28-44% over several hPSC lines as determined by flow cytometry for cardiac troponin T expression (cTnT) (Fig S1).

Cell Stimulation and Cell Surface Area

Cardiomyocytes were dissociated into single cells 20 days after the start of differentiation using TrypLE Select (Invitrogen) for 30 minutes at 37°C and plated onto Matrigel-coated coverslips (25,000 cell/cm²) in serum-free BEL medium. After 10 days, the culture medium was placed with BEL containing 100 μM PE (Sigma),100 nM ISO (Sigma), 5% serum, 20% serum (Greiner (Serum A) or Gibco (Serum B)), or left untreated for 72 h. After 72 h, all coverslips were fixed in 2% paraformaldehyde for 30 minutes. For long-term experiments, cells on coverslips were exposed to serum for 7 days followed by 72 h treatment ±PE prior to fixation. For reversibility
Figure 2) Response of hESC-CM to serum. A) Cell surface area of hESC-CM treated with 100 μM PE, 100 nM ISO, 5% serum or 20% serum for 72 h compared to untreated n≥72 cells. * = p<0.05 untreated vs. treated. B) Cell surface area of hESC-CM treated with 5% or 20% serum for 7 days 100μM PE for 72h.*=p<0.05 no PE added vs. PE-treated, n≥40. C) Representative immunofluorescent images of hESC-CM treated with 100 μM PE, 100 nM ISO, 5% serum or 20% serum for 72 h (short-term exposure) or 5% serum and 20% serum for 7 days (long-term exposure). Stacked bar graph of the percentage of cells with sarcomeric striations (visible well-organized striations (+), some (disorganized) striations (+/-), and poor/non-existent striations (-)); n≥72 cells. Red: α-actinin, Blue: DAPI, scale bar: 10 μm. D) Relative change in beating frequency of hESC-CM after immediate addition of 100 μM PE, 100 nM ISO, 5% serum, 20% serum or vehicle only control. Results normalized to initial beating frequency. * =p<0.05 vehicle only vs. treated, n≥7 measurements. E) Relative change in beating frequency of long-term serum treated hESC-CM after immediate addition of 100 μM PE, 100 nM ISO or vehicle only control. Results normalized to initial beating frequency. *=p<0.05 vehicle only vs. treated, n≥5 measurements.
experiments, cells were treated as above and subsequently refreshed with serum-free BEL medium for 3, 5, or 10 days.

For determination of cardiomyocyte cell surface area, fixed cells were labeled with anti-α-actinin (1:800; Sigma) followed by Cy3 (ImmunoResearch) secondary as previously described [32]. Nuclei were labeled with DAPI. For cell surface area measurements, 9 to 15 areas were randomly selected from each 10 mm coverslip and visualized using a Nikon Eclipse Ti-S microscope. Single plane of 1200x1600 pixels images at 313nm/pixel resolution were recorded using a Plan-fluor 20x/0.50 lens and analyzed using ImageJ software. Representative sarcomere images were visualized using a Leica TCS SP5 confocal microscope; single planes of 1024x1024 pixels images at 44nm/pixel resolution were recorded using a Plan-Apochromat 40x/1.25 oil and 100x/1.4 oil lens. Sarcomeric structures of cardiomyocytes were divided into three categories, visible well-organized striations (+; category 1), some striations mostly disorganized (+/-; category 2) and poor or non-existent striations (-; category 3).

Calcium imaging
Cell cultures undergoing cardiac differentiation were dissociated after 20 days using TrypLE Select and replated on 96-well Matrigel-coated imaging plates (BD Bioscience, 40,000 cells/cm2). In order to determine calcium transient/contraction frequency 13 days over plating, cells were loaded with 5 μM fluo-4am (Invitrogen) for 15 minutes in the presence of 0.2% Pluronic in Tyrode’s solution (140 mM NaCl, 10 mM glucose, 5 mM HEPES, 5.4mM KCl, 1.2 mM MgCl2, 1.8 mM CaCl2, pH 7.4). The cells were then washed 3 times with Tyrode’s solution and maintained in 100μl of Tyrode’s solution. With the plate maintained at 37ºC, time-lapse of Fluo-4am fluorescence was recorded using a Leica AF6000 microscope equipped with a Hamamatsu EM-CCD camera. Contracting areas were identified and imaged consecutively for 1.4 minutes (1500 images at 51ms per cycle). After recording a baseline for 20 seconds drug or serum additions were made to achieve final concentrations of 100 nM ISO, 100 μM PE, 5% serum, 20% serum, or vehicle only. Calcium transient frequency before and after treatment was determined using ImageJ and Z-profile_ImJ software, tailored for use here to convert raw data to dF/F results [33].

Statistical Analysis
Results are expressed as mean±SEM. Comparisons were made using one-way ANOVA with Bonferroni’s multiple comparison post test. Values of p ≤0.05 were considered significant. Statistical analyses were performed using GraphPad Prism.

Results
A hypertrophic response to serum in neonatal rat cardiomyocytes
We first confirmed well established hypertrophic effects of serum and PE on Rat-CM using cell surface area and beating frequency as standard surrogate measures. Rat-CM were plated in the original isolation medium containing 20% serum or in the absence of serum (Fig 1A, 0h). After 12 h, the cells were transferred to 20, 5 or 0% serum and maintained at this serum concentration until the end of the experiment (60h). Although the initial plating of these cells with or without serum (12 h exposure) did not affect cell surface area (Fig 1B), by 36h, cells in serum (5% or 20%) were significantly larger than those in absence of serum (Fig 1B). By 60h, a concentration-dependent effect was evident with cells maintained in 20% serum now significantly the largest (Fig 1C) there was no observed difference in the sarcomeric structures or organization between the groups (Fig S2).

Exposure of cardiomyocytes to PE has been previously shown to increase their cell surface area [34]. We observed a similar hypertrophic effect with exposure to PE for cells maintained in 0% or 5% but not 20% serum (Fig 1C). It was further noted that Rat-CM beat significantly more slowly in the absence than in the presence of serum (Fig 1D, n≥7, p<0.05). PE exerted an expected positive chronotropic action under serum-free conditions (Fig 1D, n≥7, p<0.05), effects which were similar to serum. Cells maintained in serum were refractory to the positive chronotropic effect of PE.
Figure 3) The stability of serum-induced alterations in hESC-CM. A) Schematic of treatment protocol. Cell surface area of hESC-CM treated with 5% or 20% serum for 3 days (n≥81) (B) or 7 days (n≥42) (C) and then removed for 3, 5, or 10 days. *p<0.05 serum vs. removal. D) Representative immunofluorescent images of hESC-CM treated with 100 μM PE, 5%, and 20% serum for 72 h +3 days (short-term exposure +3 days) or 5% or 20% serum for 7 days +3 days (long-term exposure +3 days) and then removed for 3 days (short-term exposure-3d and long-term exposure-3d, respectively). Stacked bar graph of the percentage of cells of different sarcomeric structural class (visible well-organized striations (+; category 1), some (disorganized) striations (+/-; category 2), and poor/non-existent striations (-; category 3)); n≥72. Red: α-actinin, Blue: DAPI, scale bar: 10μm.
A hypertrophic and positive chronotropic response of hESC-CM to serum containing medium
We assessed whether hESC-CM responded similarly to Rat-CM on exposure to serum or known
adrenergic stimuli. When measured after 72h, cells maintained in 5% or 20% serum (2 batches,
Sigma (Serum A) and Gibco (Serum B)) or PE all showed increased cell surface area (Fig 2A). The
β-adrenergic agonist ISO failed to induce a significant increase (Fig 2A). Combined exposure of
cells to both serum (>7days) and PE (72h) did not induce any additional increase in cell surface
area (Fig 2B).

As results for serum brands A and B were similar, only serum B was used in subsequent
experiments. Interestingly, besides the increased cell surface area, hESC-CM exposed to 5% or
20% serum developed abnormal sarcomeric structures. These effects were quantified and cells
were divided into three categories: 1) visible well-organized striations, 2) some (disorganized)
striations, and 3) poor/non-existent striations. The percentage of cells in category 1 was reduced
and in category 3 was increased (Fig 2C) as early as 72 h after serum addition and was further
exacerbated by long term serum exposure (Fig 2C).

Stimulation of the α-adrenergic system with PE or the β-adrenergic system with ISO has been
shown previously to induce a positive chronotropic response in hESC-derived cardiomyocytes
[3]. We confirmed these responses in this model. 100 μM PE induced a 2.25 fold relative increase
in beating frequency (n=11 beating clusters) and 100 nM ISO induced a 1.9 fold relative increase
(n=11 beating clusters) (Fig 2D). Furthermore, we found that serum addition to a serum-free
culture caused a similar increase in beating frequency. Following chronic exposure to serum (>7
days) the positive chronotropic effect of PE was still evident but the effect of ISO was completely
lost (Fig 2E). Besides the changed in beating frequency all other measurement parameters of the
calcium transient (e.g. upstroke, slope decay and amplitude) showed no significant difference
between treated and untreated groups (data not shown).

The hypertrophic effect of serum on hESC-CM may be irreversible.
To assess the stability of the serum- or PE-induced hypertrophy we measured cardiomyocyte
surface area upon withdrawal of these factors (Fig. 3A). For cells exposed to serum or PE for 72
h, there was no significant decrease in cell surface area at any time point and the cells remained
significantly larger than the untreated cells (Fig 3C). However cells treated with serum for 7 days
and then deprived of serum showed a significant decrease in cell surface after 5 days in serum-
free medium (Fig 3B). By this time point these cells were no longer significantly larger than the
untreated cells.

The sarcomeric structures which had already remodeled in the presence of serum appeared
even more disrupted, with many more cells showing disorganized or missing striations (Fig 3D).
This loss of sarcomeric structure was increased the longer the cells were deprived of serum (data
not shown). The removal of PE from the cells had little or no effect on the sarcomeric structure (Fig 3D).

Serum effects on cardiac hypertrophic disease modeling.
Since hiPSC-CM from patients with gene mutations affecting the heart are now being increasingly
used to model cardiac disease, it is important to determine whether serum, still widely used
in the culture of cardiomyocytes, would affect the measurement of the disease phenotype in
culture. To address this, we used a series of control and disease hiPSC lines recently derived
in our laboratory (Chapter 6), differentiated them to cardiomyocytes and assessed the impact
of exposure to serum or adrenergic agonists on cell surface area and automatcity. hiPSC-CM
derived from a healthy individual and three patients with a mutation in the MYBPC3 gene linked
to HCM (Chapter 6) were kept in serum-free medium and were treated with 5% or 20% serum
or left untreated for 3 or 7 days. After 3 days, HCM-iPSC-CM kept serum-free were significantly
larger than the ctrl-hiPSC-CM with a ≥1.5 fold increase in cell surface area (Fig 4A). The healthy
control hiPSC-CM displayed a 3-fold increase in surface area on exposure to either 5% or 20%
serum (Fig 4B and C). The hiPSC-CM from the HCM disease patients were significantly larger than
the control in serum-free conditions, suggesting that they had become hypertrophic without additional stimuli, but these cells failed to show a further increase in surface area in response to serum (Fig 4A-C). Similarly to the hESC-CM under serum-free conditions, the surface area of the control hiPSC-CM increased in response to a 72h exposure to 100μM PE (Fig 4D and 100nM of ISO showed no effect (Fig S3)), whereas the hiPSC-CM from the three disease patient lines showed no response (Fig 4D and Fig S3).

Longer-term (>7 days) exposure of hiPSC-CM to serum produced similar results as short-term exposure (72 h): the healthy control hiPSC-CM showed an increase in cell surface area while the HCM-hiPSC-CM showed no change upon serum exposure (Fig 4E). Reflecting the hESC-CM results in Fig 2B, hiPSC-CM also failed to show an additional hypertrophic response to PE when maintained in serum (Fig 4E).

Upon examination of the sarcomeric structures of the hiPSC-CM the percentage of cells in category 3 was increased after 72 h treatment with serum only in the healthy control hiPSC-CM while the percentage of category 3 cells remained relatively the same in the HCM-hiPSC-CM (Fig 4F-H).

Discussion

There are numerous reports documenting the adverse effects of serum on the efficiency of cardiac differentiation of PSCs [35;36]. Nevertheless serum is often used as a standard medium additive to improve cardiomyocyte viability both during and after differentiation by providing growth factors, nutrients and hormones [19]. However, the effect of serum on cardiomyocytes during their long-term maintenance in culture has not been investigated in much depth even though several studies have described hypertrophic effects of serum on primary rodent cardiomyocytes in culture [37;38]. For this reason, serum concentrations are usually reduced or absent in experiments carried out on these cells. In this study we first confirmed the published effects of serum and PE on primary neonatal Rat-CM and then investigated the effects of serum on cardiomyocytes derived from hESC and hiPSC, both from a healthy individual and patients with a mutation in the MYBPC3 gene causing HCM. We found that the hypertrophic and physiological effects on the Rat-CM were as reported but the effects on the hPSC derived cardiomyocytes varied depending on how long they were exposed to serum (72h or >7 days) and which concentration had been present. Hypertrophic and physiological effects of serum were only seen in healthy hPSC and not in cardiomyocytes from the hiPSC lines derived from patients carrying the HCM associated mutation.

hESC-CM and hiPSC-CM derived from the healthy individual showed similar responses to serum, with an increased cell surface area and altered beat rate. They were also similar in their responses to PE and ISO. However the hiPSC-CM derived from patients with a mutation causing HCM showed no response to the addition of serum; they were larger than controls at the outset in serum free conditions and this did not increase further. These cells were also unresponsive to PE and ISO. The underlying mechanism for this is not clear but is being investigated in an independent study.

In the hESC-CM and healthy hiPSC-CM, short-term serum (72 h) exposure resulted in similar changes in cell surface area and beating frequency as PE, a known adrenergic agonist and inducer of hypertrophy in cardiomyocytes in vitro. This effect was not reversed by the removal of serum, even after 10 days. In addition, ISO had no effect on the beating frequency of hPSC-CM exposed to serum while in the absence of serum cells doubled their beating frequency as expected (Fig 2D and 2E). Our results suggest serum may control cardiac hypertrophy through the adrenergic pathway while additional unknown components in serum, and not present in the PE or ISO conditions, could cause the adverse effects on the sarcomeric structure. In concordance, previous studies demonstrated that isolating and maintaining Rat-CM in serum-free medium resulted in a decrease in stress-like fibers which are more commonly seen in cardiac non-muscle cells [21]. Moreover, studies in isolated human and chick cardiomyocytes have demonstrated the presence of stress-fibers both in unhealthy [39] and immature [40] cardiomyocytes. On the
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On the other hand, it is possible that the breakdown of sarcomere structure may only be remodeling of cell structure due to chronotropic or inotropic changes caused by unknown components in the serum. However, the complexity of the adrenergic pathway and the undefined factors in serum require additional investigation. Previous studies on Rat-CM have revealed various parts of the hypertrophy pathway with differing responses to serum [37;38;41]. In one report, the endothelin 1 hypertrophic pathway was suppressed by retinoic acid but serum-induced hypertrophy was not suppressed [38]. In a later study PE and serum-induced hypertrophy were both found to be impaired by the overexpression of CHAMP [41]. Taken together with our results and the lack of an effect of ISO on the beating frequency of hESC-CM after long-term exposure to serum, it might be inferred that serum affects the same pathway as β-adrenergic stimuli rather than α-adrenergic stimuli.

When hESC-CM or healthy hiPSC-CM were treated with serum for longer periods of time (> 7 days) they also showed an increase in cell surface area but lost their augmented response to PE. While other drugs were not tested, these results implied that serum in cell cultures could mask, or otherwise alter, drug induced effects. Ren et al (2012) for example showed glucocorticoids induced hypertrophy in rat embryonic cardiomyocytes (H9C2 cells) cultured in serum but in the absence of serum, glucocorticoids protected the cardiomyocytes from apoptosis. Furthermore, this increased cell surface area reverted when the serum was removed for at least 5 days [23. However the exact cause of this phenomenon is unclear. It may be that the sudden withdrawal of 

Figure 4) Response of hiPSC-CM to serum. Cell surface area of hiPSC-CM derived from a healthy individual (healthy) and three patients with hypertrophic cardiomyopathy (HCM1, HCM2, HCM3) in (A) serum-free medium *p=0.05 healthy vs. HCM n≥60 cells. (B) treated with 5% for 72 h. *p=0.05 healthy vs. HCM, n≥66 cells. (C) treated with 20% serum for 72 h. *p=0.05 healthy vs. HCM, n≥64 cells. D) Cell surface area of hiPSC-CM ± 100 μM PE for 72 h. *p=0.05 no PE added vs. PE-treated, n≥50 cells. E) Cell surface area of hiPSC-CM treated with 5% or 20% serum for 7 days ± 100μM PE for 72h, n≥39 cells. F-H) Stacked bar graph of the percentage of cells of different sarcomeric structural class (visible well-organized striations (+; category 1), some (disorganized) striations (+/-; category 2), and poor/ non-existent striations (-; category 3)); F: no serum; G: treated with 5% serum; H: treated with 20% serum; n≥47.
of growth factors from the serum leads to problems in basic cell metabolism and the synthesis and trafficking of sarcomeric proteins has become comprised. Thus the cardiomyocytes with longer exposure to serum may be on their way to losing viability. Others have previously reported similar results in Rat-CM in which serum-deprivation lead to increased apoptosis [42].

In addition to PSC-CM from healthy individuals, we investigated the effects of serum on cells derived from three patients with HCM caused by a mutation in MYBPC3 gene. Clinically these patients had an increase interventricular septum thickness but other heart functions, such as fractional shortening, left ventricular systolic-end and diastolic-end diameter, remained within the normal range [43]. Their derivative hiPSC-CM were resistant to PE and ISO induced hypertrophy and chronotropic effects and they also lacked the response to serum seen in the controls and primary cardiomyocytes. Thus their enlarged cell surface areas and altered beat rates relative to controls in the absence of serum were no longer evident in the presence of serum. This represents a cautionary note on the culture conditions used to compare diseased and control cardiomyocytes derived from pluripotent stem cells. In addition, some experiments described in the literature use primary rodent cells ±24 h after transfer to serum-free medium but this may not always be the case. Improved consistency in the results may be achieved by carrying out the experiments under serum-free or low serum conditions.

In conclusion, we have demonstrated that the addition of serum to PSC-CM derived under defined and conventional serum-free conditions can significantly alter the phenotype of cardiomyocytes: their surface area, sarcomeric structure and beating frequency are all parameters that show relevant alterations. These are among the parameters widely used to report phenotypes in cardiomyocytes derived from patient hiPSC. Therefore it is highly desirable to control experimental conditions and to culture hPSC derived cardiomyocytes in fully defined culture media.

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References

1. Kehat, I., Gepstein, A., Spira, A. et al (2002) High-resolution electrophysiological assessment of human embryonic stem cell-derived cardiomyocytes: a novel in vitro model for the study of conduction Circ.Res. 91, 659-661
2. Mummery, C., Ward, D., van den Brink, C.E. et al (2002) Cardiomyocyte differentiation of mouse and human embryonic stem cells J.Anat. 200, 233-242
3. Mummery, C., Ward-van, O.D., Døevendans, P.et al (2003) Differentiation of human embryonic stem cells to cardiomyocytes: role of coculture with visceral endoderm-like cells Circulation 107, 2733-2740
4. Braam, S.R., Tertoolen, L., van de Stolpe, A.et al (2010) Prediction of drug-induced cardiotoxicity using human embryonic stem cell-derived cardiomyocytes Stem Cell Res. 4, 107-116
5. Braam, S.R., Tertoolen, L., Casini, S.et al (2013) Repolarization reserve determines drug responses in human pluripotent stem cell derived cardiomyocytes Stem Cell Res. 10, 48-56
6. He, J.Q., January, C.T., Thomson, J.A.et al (2007) Human embryonic stem cell-derived cardiomyocytes: drug discovery and safety pharmacology Expert.Opin.Drug Discov. 2, 739-753
7. Liang, P., Lan, F., Lee, A.et al (2013) Abnormal Calcium Handling Properties Underlie Familial Hypertrophic Cardiomyopathy Pathology in Patients-Specific Induced Pluripotent Stem Cells Cell Stem Cell 12, 101-113
8. Moretti, A., Bellin, M., Welling, A.et al (2010) Patient-Specific Induced Pluripotent Stem-Cell Models for Long-QT Syndrome N.Engl.J.Med.
9. Carvajal-Vergara, X., Sevilla, A., D’Souza, S.L.et al (2010) Patient-specific induced pluripotent stem-cell-derived models of LEOPARD syndrome Nature 465, 808-812
10. Burridge, P.W., Thompson, S., Millrod, M.A.et al (2011) A universal system for highly efficient cardiac differentiation of human induced pluripotent stem cells that eliminates interline variability PLoS.One. 6, e18293
11. Ng, E.S., Davis, R., Stanley, E.G.et al (2008) A protocol describing the use of a recombinant protein-based, animal product-free medium (APEL) for human embryonic stem cell differentiation as spin embryoid bodies Nat.Protoc. 3, 768-776
12. Ren, Y., Lee, M.Y., Schliffke, S.et al (2011) Small molecule Wnt inhibitors enhance the efficiency of BMP-4-directed cardiac differentiation of human pluripotent stem cells J.Mol.Cell Cardiol. 51, 280-287
13. Mummery, C.L., Zhang, J., Ng, E.S.et al (2012) Differentiation of human embryonic stem cells and induced pluripotent stem cells to cardiomyocytes: a methods overview Circulation Research 111, 344-358
14. van der Valk, J., Brunner, D., De Smet, K.et al (2010) Optimization of chemically defined cell culture media–Replacing fetal bovine serum in mammalian in vitro methods Toxicology in Vitro 24, 1053-1063
15. Bass, G.T., Ryall, K.A., and Katikapalli, A. (2012) Automated image analysis identifies signaling pathways regulating distinct signatures of cardiac myocyte hypertrophy J.Mol.Cell Cardiol. 52, 923-930
16. Nag, A.C., Lee, M.L., and Kosiur, J.R. (1990) Adult cardiac muscle cells in long-term serum-free culture: myofibrillar organization and expression of myosin heavy chain isoforms In Vitro Cellular & Developmental Biology

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22 Peters, M.F. and Scott, C.W. (2012) Evaluation of Cellular Impedance Measures of Cardiomyocyte Cultures for Drug Screening Applications Assay Drug Dev Technol. 10, 525-533
23 Ren, R., Oakley, R.H., Cruz-Topete, D.et al (2012) Dual role for glucocorticoids in cardiomyocyte hypertrophy and apoptosis Endocrinology 153, 5346-5360
24 Simpson, P., McGrath, a., and Savion, S. (1982) Myocyte hypertrophy in neonatal rat heart cultures and its regulation by serum and by catecholamines Circulation Research 51, 787-801
25 Birket, M.J., Casini, S., Kosmidis, G.et al (2013) PGC-1alpha and Reactive Oxygen Species Regulate Human Embryonic Stem Cell-Derived Cardiomyocyte Function Stem Cell Reports 1, 560-574
26 Yu, M., Xiang, F., Beyer, R.P.et al (2010) Transcription Factor CHF1/Hey2 Regulates Specific Pathways in Serum Stimulated Primary Cardiac Myocytes: Implications for Cardiac Hypertrophy Current genomics 11, 287-296
27 Pijnappels DA, Schalij MJ, van TJ et al. (2006) Progressive increase in conducportion velocity across human mesenchymal stem cells is mediated by enhanced electrical coupling Cardiovasc.Res. 97, 171-181
28 Elliott, D.A., Braam, S.R., Koutsis, K.et al (2011) NKX2-5(eGFP/w) hESCs for isolation of human cardiac progenitors and cardiomyocytes Nat.Methods 8, 1037-1040
29 Dambrot, C., Van Den Berg, C., Oostwaard, D.W., Davis, R., Braam, S., Ng, E., and Mummery, C. (2012) In Human Stem Cell Manual (Second Edition) pp. 413-431, Academic Press, Boston
30 Dambrot, C., van de Pas, S., van, Z.L.et al (2013) Polycistronic lentivirus induced pluripotent stem cells from skin biopsies after long term storage, blood outgrowth endothelial cells and cells from milk teeth Differentiation 85, 101-109
31 Goncalves, M.A., Janssen, J.M., Holkers, M.et al (2010) Rapid and sensitive lentivirus vector-based conditional gene expression assay to monitor and quantify cell fusion activity PLoS One 5, e10954
32 Davis, R.P., Casini, S., van den Berg, C.W.et al (2012) Cardiomyocytes derived from pluripotent stem cells recapitulate electrophysiological characteristics of an overlap syndrome of cardiac sodium channel disease Circulation 125, 3079-3091
33 Takahashi, A., Camacho, P., Lechleiter, J.D.et al (1999) Measurement of intracellular calcium Physiol Rev. 79, 1089-1125
34 Umar, S., van der Valk, E.J., Schalij, M.J.et al (2009) Integrin stimulation-induced hypertrophy in neonatal rat cardiomyocytes is NO-dependent Mol.Cell Biochem. 320, 75-84
35 Freund, C., Ward-van, O.D., Monshouwer-Kloots, J.et al (2008) Insulin redirects differentiation from cardiogenic mesoderm and endoderm to neuroectoderm in differentiating human embryonic stem cells Stem Cells 26, 724-733
36 Passier, R., Oostwaard, D.W., Snapper, J.et al (2005) Increased cardiomyocyte differentiation from human embryonic stem cells in serum-free cultures Stem Cells 23, 772-780
37 Akazawa, H. and Komuro, I. (2003) Roles of cardiac transcription factors in cardiac hypertrophy Circulation Research 92, 1079-1088
38 Zhou, M.D., Sucov, H.M., Evans, R.M.et al (1995) Retinoid-dependent pathways suppress myocardial cell hypertrophy Circulation Research 79, 2268-2278
39 Bird, S.D., Doevendans, P.A., van Rooijen, M.A.et al (2003) The human adult cardiomyocyte phenotype Cardiovasc.Res. 58, 423-434
40 Dlugosz, a.a., Antin, P.B., Nachmias, V.T.et al (1984) The relationship between stress fiber-like structures and nascent myofibrils in cultured cardiac myocytes The Journal of cell biology 99, 2268-2278
41 Liu, Z.P. and Olson, E.N. (2002) Suppression of proliferation and cardiomyocyte hypertrophy by CHAMP, a cardiac-specific RNA helicase Proceedings of the National Academy of Sciences of the United States of America 99, 7391-7395
42 Zhu, H., McElwee-Witmer, S., Perrone, M.et al (2000) Phenylephrine protects neonatal rat cardiomyocytes from hypoxia and serum deprivation-induced apoptosis Cell death and differentiation 7, 773-784
43 van Dijk, S.J., Dooijes, D., dos, R.C.et al (2009) Cardiac myosin-binding protein C mutations and hypertrophic cardiomyopathy: haploinsufficiency, deranged phosphorylation, and cardiomyocyte dysfunction Circulation 119, 1473-1483

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Supplementary Figures:

Fig S1) Cardiac differentiation efficiency of the Nkx 2-5eGFP/w hESC line and LUMC0004ictrl determined by flow cytometry of cardiac troponin T (cTnT) expression (n=11 for Nkx 2-5eGFP/w, two independent experiments for LUMC0004ictrl).

Fig S2) Stacked bar graph of the percentage of RAT-CM with sarcomeric structural class (visible well-organized striations (+; category 1), some (disorganized) striations (+/-; category 2), and poor/non-existent striations (-; category 3)); n≥41 cells.

Fig S3: Response of hiPSC-CM to isoproterenol (ISO). Cell surface area of hiPSC-CM ± 100 nM ISO for 72 h (n≥25 cells).
