Fluorescence spectroscopy of low-level endogenous β-adrenergic receptor expression at the plasma membrane of differentiating human iPSC-derived cardiomyocytes

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Abstract

The potential of human induced pluripotent stem cells (hiPSCs) to be differentiated into cardiomyocytes (CMs) mimicking the adult CMs functional morphology, marker genes and signaling characteristics has been investigated since over a decade. The evolution of the membrane localization of CM-specific G protein-coupled receptors throughout differentiation has received, however, only limited attention to date. We employ here advanced fluorescent spectroscopy, namely linescan Fluorescence Correlation Spectroscopy (FCS), to observe how the plasma membrane abundance of the β₁- and β₂-adrenergic receptors (β₁/₂-ARs), labelled using a bright and photostable fluorescent antagonist, evolves during long-term monolayer culture of hiPSC-derived CMs. We compare it to the kinetic of observed mRNA levels in wildtype (WT) hiPSCs and in two CRISPR/Cas9 knock-in clones. We conduct these observations against the backdrop of our recent report that β₂-ARs, as opposed to β₁-ARs, specifically segregate to the T-Tubular system of adult CMs.

Introduction

Human induced pluripotent stem cells (hiPSCs) can be differentiated into cardiomyocytes (CMs) that are functionally comparable to embryonic stem (ES) cell-derived CMs and/or mimicking the functional morphology of adult cardiomyocytes. Although fully differentiated hiPSCs do not yet achieve a phenotype entirely reminiscent of that of adult CMs - an area of active and intense research [1] - the possibility of driving in a controlled way these cells from a pluripotent towards a more mature
phenotype offers several important unique opportunities. First and foremost, undifferentiated hiPSCs are amenable to genetic editing, which allows them to incorporate knockouts of selected genes or addition of functional tags, such as fluorophores, that will then be carried forward to the CM phenotype.

Secondly, expression of key genes changes over time and this can be followed in detail, providing an insight into how key structures of the CM phenotype originate, and how specific signaling pathways and their associated molecular mechanisms develop. In this respect, several works have explored since about 10 years the evolution of the transcriptome of hiPSCs as they directionally differentiate towards CMs. The majority of CM-specific genes seem to be upregulated after day 14 [2] [3]. More recently, single cell sequencing has allowed to shed light also on cell-to-cell expression variability for selected genes [4] [5].

An important feature of adult CMs is the ability to increase or reduce both strength and frequency of their contractions in response to external stimuli, two processes that go under the name of inotropic and chronotropic responses, respectively. The β-adrenergic system, embodied by the β₁- and β₂-adrenergic receptors (β-ARs, expressed from the ADRB1 and ADRB2 genes respectively), represents the typical pathway mediating catecholamine-driven inotropic and chronotropic responses, and is a crucial therapeutic target in the context of heart failure, mostly thanks to the discovery of the beneficial action of negative inotropes/chronotropes such as β-blockers. In the adult heart, β₁ and β₂ are both expressed, with a ratio of β₁/β₂-ARs of around 70%–80%/30%–20% in human ventricles [6-8].

The evolution of endogenous β-AR expression over maturation time in hiPSCs directionally differentiated towards CMs has thus been explored in several recent works. Wu et al. observed mRNA levels and effects on contractility of β-AR expression in differentiating hiPSC, in a model of dilated cardiomyopathy [6]. Jung et al. studied adrenoreceptor mRNA levels within the context of functional remodelling [9], while Hasan et al. monitored how receptor-specific cAMP signaling evolves throughout hiPSC differentiation towards CMs [10]. Kondrashov et al. studied the temporal dynamics of β₂-AR expression in CRISPRed hiPSC [11]. An earlier work by Yan et al. studied the effect of adrenergic signalling on regulating cardiac differentiation of mouse embryonic stem cells and monitored ADRB1 and ADRB2 expression as a function of time [12].

These works interestingly report a predominant ADRB2 expression with respect to ADRB1, resulting in a higher second messenger production upon selective β₂-AR stimulation, that then decays during differentiation. This results in a β₂-AR dominated functional response (contractility) which is then caught-up by the β₁-AR at day 60. These results point to the interesting and persistent gene expression dynamics affecting the relative weight of these two key adrenoreceptors. However, transcriptomic analyses and functional downstream readouts, even if conducted at the single cell level, fail to provide information about what occurs in terms of 1. protein expression and 2. receptor
localization. We believe this is an important aspect to consider, in the light of our recent observation of the peculiar differential localization of the $\beta_1$- and $\beta_2$-ARs on the membranes of adult cardiomyocytes [13]. We could show that the $\beta_2$-AR partitions to the T-tubular network of the cardiomyocyte and does not appear to diffuse upon the basolateral (‘crest’) surface plasma membrane of the cell, thus explaining the compartmentalized cAMP signal observed earlier in this cell-type [14].

Key to our observation was the use of a high-sensitivity fluorescence spectroscopy method, namely linescan-FCS [15], that allowed us appreciating minute amounts of endogenous receptors diffusing on the membranes using a fluorescent antagonist (JE1319). In that context we could also observe, not only significant CM to CM variability of receptor expression (i.e. $\beta_1$-AR is visible only in 60% of the cells considered), but also the fact that, in CM-like cells such as H9c2 and CM-differentiated hiPSC, $\beta_2$-ARs were clearly seen diffusing on the basolateral membrane.

We thus reasoned that it is now timely to investigate how the plasma membrane concentration of the $\beta_1$- and $\beta_2$-adrenergic receptors evolves during long-term monolayer culture of hiPSC-CM, whilst comparing it to the measured mRNA levels. Moreover, considering that one of the key advantages of hiPSCs is their ability of being gene-edited in their pluripotent, non-differentiated state, we extended our analysis also to two hiPSC clones carrying a heterozygous and a homozygous CRISPR/Cas9-mediated knock-in of a fluorescent tag at the N-terminus of the $ADRB2$. This should further allow to corroborate the missing link between protein localization and mRNA levels.

We could observe that in wildtype (WT) hiPSCs, there is a discrepancy between $\beta_2$-AR appearance at the surface plasma membrane during early maturation stages (day25) with respect to the measured mRNA levels, and a significant cell-to-cell heterogeneity in receptor abundance at the plasma membrane. At later maturation stages (day60) mRNA levels correlated with membrane appearance of $\beta_2$-AR in WT hiPSC-CM as well as in CRISPRRed hiPSC-CM clones, indicating that the addition of a tag at the N-terminus of $ADRB2/\beta_2$-AR still allowed for correct receptor transcription, expression and localization at and after day 60.

These observations open fascinating questions on the mechanisms that control and time receptors appearance and distribution at the plasma membrane, with potential implication towards better understanding of a signaling cascade at the core of heart pathophysiology and failure.

**Results**

We first set to establish a CM-differentiation assay for hiPSCs (See Methods), identifying timepoints (0, 25, 60 and 100 days) after onset of differentiation for our
investigation (Supplementary Figure 1). The hiPSCs morphology was investigated by transmitted light microscopy, as reported in Figure 1a and Supplementary Figure 1a.

We worked with three hiPSC lines - WT hiPSCs, and two clonal lines arising respectively from a heterozygous (66-35) and homozygous knock-in (16-31) of the monomeric Enhanced Green Fluorescent Protein (mEGFP) at the N-terminus of the endogenous ADRB2 gene locus. Clones 16-31 and 66-35, created using CRISPR/Cas9 editing in combination with a novel partially single-stranded, DNA-based hybrid HDR donor cocktail, and details of the knock-in procedure as well as the clones’ identification are described in Supplementary Figure 2 and the Methods.

Figure 1a further displays fluorescence excitation of the cells at 488 nm, indicating a substantial amount of autofluorescence [16] [17], that increases as cells differentiate towards a mature CM phenotype. Interestingly, the level of green signal is comparable between the WT cells and the two mEGFP knock-in clones, suggesting that the very low-level expression of the tagged β2-AR has to compete with autofluorescence, and that alternative tagging strategies shall be considered [18, 19]. In the following, we therefore employed the fluorescent antagonist which we successfully used in our recent spectroscopic imaging of adrenergic receptors in adult murine cardiomyocytes (JE1319 [13]), while still exploiting the information arising from the two knock-in clones.

Figure 1b displays the fully differentiated hiPSC-CM phenotype at day ~100, employing α-actinin and cardiac troponin T immunofluorescence to highlight the presence of sarcomeric structures and thus the success of differentiation. This shall be compared to the lower immunofluorescence signal observed in the few non-CM cells present in the culture (dashed lines in the DIC and Merge overlay of Fig. 1B). Morphologically, there are no detectable differences between the WT cells and clones 16-31 and 66-35.

When subjected to selective adrenergic stimulation (for either β1-AR using ICI-118551 or β2-AR using CGP-20712A) (Figure 1c), the day ~100 hiPSC-CM displayed a chronotropic response, as measured by their beating rate, comparable between the WT cells and the homozygous clone 16-31. In particular, the selective β2-AR stimulation displayed an increase in beating rate almost identical between the WT and the knock-in, suggesting that N-terminal tagging of the β2-AR is neither affecting its proper expression nor its function. This is also displayed in Videos S1 and S2, which show spontaneously contracting monolayers of the day 100 clone 16-31 before and after β2-AR stimulation, respectively. These results also confirm that, almost undetectable levels of expressed receptor can contribute to the normal response of the cell to external stimuli [13].

Having established and characterized our three hiPSC-CM cell lines, we now set to explore in detail the kinetic of β2-ARs expression as a function of the maturation time. In order to do so, we first performed quantitative mRNA measurements using qRT-PCR (see Methods). Here, the presence of a tag in the two knock-in clones further allows to measure the expression levels of individual alleles, as well as to compare transcript levels from a tagged variant of the β2-AR to its WT expression. The results of this
experiment are displayed in Figure 2: here, we compare absolute mRNA levels (Figure 2a) extracted from cultures of WT hiPSC and our two clones. Two sets of qPCR primers, as detailed in the methods, one complementary to the mEGFP sequence (green bars) and the other complementary to the ADRB2 nucleotide sequence (black bars) were used, thus allowing to compare expression level of the endogenous background with respect to the tagged allele(s) (see Methods).

Figure 1 Characterization of hiPSC-CMs. (A) Comparison of undifferentiated (day 0) and differentiated (day 100) WT hiPSC-CM and mEGFP-ADRB2 CRISPRRed clones 16-31 and 66-35. Fluorescence images represent autofluorescence in the green spectral range at 488 nm. Shown are representative images of 2 replicates with imaging of ≥4 points-of-view. Contrast settings are identical. Scale bar is 50 µm. (B) Immunofluorescence staining of day 100 WT hiPSC-CM and clones 16-31 and 66-35 for typical cardiomyocyte markers α-actinin and cardiac troponin T (cTnT). Nuclei were stained with NucBlue. Insets show magnifications of regions marked with white rectangles in DIC. Dashed lines in DIC and merge indicate cells lacking a
cardiac phenotype, most likely fibroblasts. Representative images of 3 replicates with imaging of ≥3 points-of-view are shown. Scale bar is 50 µm and in magnification 5 µm. (C) Functionality recording of WT hiPSC-CMs and clone 16-31 after 100 days of differentiation. Shown is the increase in beating frequency normalized to basal frequency upon stimulation with 1 µM isoproterenol (ISO) of β₁-ARs and β₂-ARs. The frequency remains unchanged upon stimulation after blocking the receptors with 10 µM propranolol (Control). Graph represents the mean ±SEM of ≥3 independent experiments.

Upon normalization to GAPDH and the day 0 expression levels of the homozygous clone 16-31, Figure 2b allows to fully appreciate the temporal evolution of ADRB2 transcripts amount along the differentiation pathway. In accordance to what was observed in previous reports [6], ADRB2 mRNA transcripts appear to be substantially upregulated already after day 25 from the onset of differentiation, with modest increases along the further timepoints. As expected, clone 16-31 displays comparable expression levels of mRNA containing the mEGFP sequence and of mRNA containing the ADRB2 coding sequence, indicative that both alleles carry the knock-in. Instead, clone 66-35 displays roughly half mEGFP mRNA than ADRB2 mRNA, indicative for both a monoallelic knock-in as well as of a roughly equal level of expression of both alleles throughout the differentiation. In the light of these observations, we ascribe here the lower overall ADRB2 mRNA levels observed in both clone 16-31 and clone 66-35 to clonal variability.

Figure 2 Indication of endogenous ADRB2 and mEGFP-ADRB2 mRNA expression during hiPSC-CM culture maturation. (A) Absolute cycle threshold (cₜ) values for GAPDH, ADRB2 and mEGFP-ADRB2 are shown. (B) Relative expression of data shown in A) normalized to GAPDH and calibrated to day 0 of clone 16-31. Data are mean ±SD of (3-6 replicates of) 2 independent experiments.

The next question is now if and how mRNA expression levels of the ADRB2 over time are mirrored by the β₂-AR expression levels at the plasma membrane. In order to measure the latter, we resort to the approach we already successfully used to monitor β-AR expression in adult CMs, based on linescan-FCS to monitor minute amounts of diffusing receptors tagged with our fluorescently labeled antagonist. Briefly, the
excitation volume of the confocal microscope is scanned rapidly (< 1 ms) along a line, positioned along the basolateral membrane of the cell. Fluctuations due to the diffusion of the tagged receptors are extracted by a mathematical analysis known as autocorrelation, to yield a curve termed autocorrelation function (ACF), that contains information both on the diffusivity and concentration of the tagged receptors. This function can be then displayed in a color code, and average ACF from several cells pooled in a heatmap image, that allows direct comparison between and across clones and timepoints. In general, the qualitative observation of an ACF, with decaying amplitude (from red – high correlation to blue – no correlation) indicates the presence and diffusion of labeled receptors. For this reason, the method can be used in a powerful way to estimate the presence of the β1-AR and β2-AR at the basolateral membrane in our hiPSC clones, and measure their evolution over time. The result of this analysis is displayed for the WT hiPSC-CM in Figure 3, comparing β1-AR and β2-AR expression over time. Most of the undifferentiated (day 0) cells display low to negligible level of β1-AR and β2-AR expression, with the exception of one cell where β2-AR was selectively labeled. Along these lines the day 25 cells investigated still displayed a substantial cell-to-cell heterogeneity in plasma membrane expression, with a significant amount of cells still displaying no-trace of either β1-AR (~90%) or β2-AR (~75%) at the cell membrane. This is a striking departure from the mRNA data reported in Figure 2, as the single cell plasma membrane receptor abundance data paint a significantly more complex, and heterogeneous, picture than the bulk mRNA levels. Both receptors become visible at the plasma membrane in most cells at 60 days from the onset of differentiation, and no qualitative difference is observed between day 60 and day 100. These results open an interesting paradox, since this would imply that at 25 days post differentiation the cell has substantial amounts of ADRB2 mRNA, but still no comparable levels of β2-AR expressed at the plasma membrane, pointing to either a limitation on translation of the receptor mRNA or of its correct targeting, as cells progress at different paces through the differentiation pathway.

![Figure 3](https://example.com/figure3.png)

**Figure 3** Linescan-FCS evaluation of β1-ARs (A) and β2-ARs (B) in WT hiPSC-CMs at day 0, 25, 60 and 100 of maturation. For each day a minimum of at least 4 scans of different hiPSC-CMs is evaluated. Colorscale indicates the amplitude of normalized autocorrelation (ACF) curves. Normalization was to each G(0) or highest peak of the curve. Each row represents the average correlation arising from the scan at the surface of one cell.
We further extracted the diffusion constants (found in Supplemental Figure 3A) and mean values range from $0.25 \sim 0.45 \, \mu m^2/s$. Supplemental Figure 3B gives an indication of the observed receptor concentrations, with mean values ranging from 3 – 12 particles in the effective detection volume of the confocal microscope. In our hands, $\beta_2$-AR expression appears higher than $\beta_1$-AR both at day 60 and day 100.

Figure 4 Comparison of $\beta_1$-ARs (A) and $\beta_2$-ARs (B) membrane expression in hiPSC-CMs at day 60 of maturation. (C) Shows the acquired linescan FCS data of hiPSC-CMs at day 60 of maturation pre-treated with 100 µM propranolol (Control). Each subpanel represents wildtype (WT) hiPSC-CMs (top), clone 66-35 (middle) and clone 16-31 (bottom) and evaluates 4 cells. Colorscale indicates the amplitude of normalized autocorrelation curves. Normalized was to each $G(0)$ or highest peak of the curve. Each bar represents the scan at the surface of one cell. Both receptor subtypes show a diffusional fingerprint at the surface plasma membrane of the WT cells and clones at day 60. Control cells indicate no detected diffusing receptors (blue).

We finally asked whether the knock-in of an N-terminal tag, that did not seem to affect mRNA levels, ended up influencing proper receptor localization at the plasma membrane. The construct corresponding to our knock-in modification of the ADRB2, appears to be well expressed and clearly localized to the plasma membrane when transfected as a plasmid into H9c2 cells (Supplemental Figure 4). Figure 4 displays, for day 60 (when full expression of the receptor on the plasma membrane is observed) that membrane expression levels are comparable between WT hiPSC-CM and the two knock-in clones, with the possible exception of $\beta_2$-AR membrane expression on the homozygous clone (16-31). Reassuringly, no membrane signal compatible with receptor diffusion was observed in the control, meaning that cells treated with the inverse agonist propranolol before being exposed to the fluorescent ligand did not display any ACF (Figure 4c).

The data from $\beta_2$-AR on day 60 of clone 16-31, display visible receptor presence at the membrane and dynamics, at least for three of the cells, whereas the other four cells do
not yield a signal compatible with receptors diffusing at the membrane. Taken together with the observation of the sizable β2-AR-mediated increase in contractility (Figure 1c), we believe that clone 16-31 is properly expressing the tagged β2-ARs to the plasma membrane, albeit displaying higher cell to cell variability.

Discussion

Our work has quantitatively explored the membrane expression level of the β1-AR and β2-AR in differentiating hiPSCs at four time points, i.e. in the undifferentiated pluripotent state, and then 25, 60 and 100 days after directed differentiation towards a CM state. Our work was conducted in parallel on WT hiPSCs, together with two CRISPR/Cas9 knock-in clones, incorporating respectively a homozygous and heterozygous N-terminal tag (mEGFP) at the β2-AR, allowing to monitor the effect of a tag on the transcription of the ADRB2 gene, as well as on the expression and localization of the β2-AR.

Membrane protein abundance, measured by linescan-FCS, a sensitive fluorescence spectroscopy approach, was then compared to mRNA levels obtained from qPCR. It shall be noted that antibodies raised against GPCRs have well documented shortcomings, and even when using anti-EGFP antibodies against the N-terminal tag in clones 16-31 and 66-35, we failed to detect receptor expression in Western blots (data not shown), reaffirming the necessity of a high-sensitivity method such as our linescan-FCS.

The approach we implemented allowed us to observe a significant discrepancy between mRNA levels of the β2-AR expression at the plasma membrane 25 days after differentiation, which raises interesting questions concerning translational regulation mechanisms for ADRB1/2 in differentiating hiPSC-CMs. Our work fills a gap in existing literature, which either addressed only mRNA levels, or functional readouts after day 30 of differentiation in hiPSCs. So far, only one work in mouse ES cells measured protein expression levels as a function of differentiation [12]. The relationship between mRNA levels and protein levels in mammalian cells is complex, and although a general correlation is expected, making transcriptional control the dominant mechanism regulating also protein abundance, there are several exceptions to this rule, as reviewed by Liu et al [20]. In differentiating cells, the situation is even more complex, and translational control has been indicated to play a major role, especially in the early stages of differentiation [21]. We shall note here that our method allows only to monitor receptor expression levels at the basolateral membrane, since we use a ligand that is non permeable to the plasma membrane [13], and we thus cannot rule out that the β2-AR is still translated, but does not localize to the plasma membrane. It could be that accessory proteins responsible for its trafficking or posttranslational modifications are not yet present in the differentiating cell at day 0 and day 25 [22]. It shall be further noted that the fully CM-differentiated hiPSCs, even at day ~100, do not display a
complete T-Tubular network, leaving open the question if the β2-AR, that in this work we observed on the basolateral membrane, would eventually segregate in these compartments [13]. Our observation of higher relative β2-AR expression levels in hiPSC-CMs appears to be in line with what was previously reported in at least three works [6, 9, 10]. Although, we shall mention here, that in recent measurements in hiPSC-derived engineered heart tissue, higher levels of β1-AR were observed throughout (private communication and [23]).

We shall also note here that our choice of mEGFP as an N-terminal tag for live cell imaging of the β2-AR, while well established in overexpression systems, faced significant challenges due to the high autofluorescence background of differentiated hiPSC-CMs, as displayed in Figure 1a. Nevertheless, the use of the tag was instrumental in confirming that the effect of the addition of a 700 bp sequence to the ADRB2 gene, resulting in a ~25 kDa tag on the final protein, did not perturb its transcription, its translation nor its final plasma membrane localization.

Overall, we can show here that linescan-FCS can be successfully used to monitor endogenous low level membrane receptor expression of GPCRs in differentiating hiPSC-CM lines, as recently demonstrated for single point FCS by Goulding et al. in HEK293 cells [18]. This substantially increases the possibilities of current methods available for the assessment of gene expression throughout hiPSC differentiation: not only we add to quantitative mRNA measurements also the dimension of actual protein expression levels, but furthermore we show how to measure it in a spatially resolved way within a single cell.

Materials and methods

Differentiation and culture of hiPSC-CM
In this work we used a well-established hiPSC line (BIHi005-A) of the stem cell core facility of the Max Delbrück Center for Molecular Medicine. Pluripotent cultures were grown at 37 °C with 5% CO2 and 5% O2, whereas differentiated cultures were maintained at 5% CO2 and atmospheric (21%) O2. Monolayers of hiPSCs were differentiated into hiPSC-CMs by modulating Wnt signaling according to a small molecule-based cardiac differentiation strategy [24, 25] followed by metabolic lactate selection in order to enrich for cardiac myocytes [26], as also previously described by us [13]. In short, hiPSC were cultured in essential 8 basal medium (E8 medium) (Gibco) on Geltrex- (Gibco) coated plates 4 days prior to cardiac induction. When they reached 80-90% confluency, mesodermal differentiation was induced on day 0 for 24 h by changing the medium to cardiac priming medium (RPMI 1640 (Gibco), 1x B27 supplement without insulin (Gibco) and 6 µM CHIR99021 (Sellek Chem)). On day 1, basal differentiation medium (cardiac priming medium without CHIR99021) was provided on top of the old cardiac priming medium. Two days later cardiogenesis was induced by changing the
medium to basal differentiation medium supplemented with 5 µM of the Wnt inhibitor IWR-1-endo (Sellek Chem). On day 5, basal differentiation medium was added on top of the old medium to let cells grow for another 2 days. Cells were cultured in maintenance medium (RPMI 1640, B27 with insulin (Gibco)) starting at day 7. Selection of and enrichment for cardiomyocytes was initiated on day 9 by culturing cells for 4 days in RPMI 1640 medium without glucose (Gibco) supplemented with 1x CDM3 supplement (Sigma Aldrich) and 5 mM sodium DL-lactate (Sigma Aldrich). From here, the beating hiPSC-CMs were cultured in maintenance medium which was exchanged every 3-5 days.

Culture of H9c2 cells
H9c2 (ATCC; CRL-1446) cells were maintained in Dulbecco’s modified Eagle’s medium (PAN-Biotech) supplemented with 10% (vol/vol) fetal bovine serum (FBS) (Sigma-Aldrich), 2 mM L-glutamine (PAN-Biotech), penicillin (100 U/mL; Gibco), and streptomycin (100 µg/mL; Gibco) at 37 °C and 5% CO₂. Cells were passaged and dissociated using 0.05%/ (ethylenedinitrilo)tetraacetic acid (EDTA) 0.02% in phosphate buffered saline (PBS) (PAN-Biotech).

Oligonucleotides

<table>
<thead>
<tr>
<th>Name</th>
<th>Sequence from 5’ to 3’</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>crRNA-103forw</td>
<td>CCTGCCGACAGACTGGCGGCGCAT</td>
<td>crRNA used in combination with generic tracrRNA to build the gRNA for Cas9 targeting to ADRB2 locus. Binds over start codon for knock-in of mEGFP</td>
</tr>
<tr>
<td>Donor plasmid for knock-in of mEGFP&lt;br&gt;Donor sequence (1.2 kbp) encoding for mEGFP plus 230bp homology arms flanked by HindIII (5’) and EcoRI (3’) restriction enzyme sites which were used for cloning into a pcDNA3.1(+) vector backbone. Served as template to generate the partially single-stranded hybrid HDR donor cocktail for knock-in of mEGFP at the ADRB2 locus using the following 4 primers:</td>
<td>&lt;br&gt;- mEGFP_N-term_f&lt;br&gt;- mEGFP_C-term_linker_r&lt;br&gt;- b2AR-hum_120bp_5’arm_f1&lt;br&gt;- b2AR-hum_120bp_3’arm_r1</td>
<td></td>
</tr>
<tr>
<td>crRNA-69forw</td>
<td>CTGCAGCCGCGATTGAGCGCA</td>
<td>crRNA used in combination with generic tracrRNA to build the gRNA for Cas9 targeting to mEGFP-ADRB2 locus. Binds over start codon for knock-in of HA signal sequence</td>
</tr>
<tr>
<td>ssODN for knock-in of HA</td>
<td>TGGGTCCGCGCTGAGGCGCCCCCAGCCAGTGCGCTCACCTGCCAGACTGCGC GCCATGAAGACGATCATCGCCCTGAGCTACATCTCTGCGGTATTCGGCCGTGAG CAAAGGCCAGAGAGCTTACCCGGGTGGTGCCCATCTGGTGAGCTGGAGCAAGGGCGAGGAGCTGTTCACCGGGGTGGTGCCCATCCTGGTCGAGCTGGACGGC</td>
<td>ssODN consisting of 60 bp homology arms used as HDR donor template for knock-in of HA signal sequence at the N-terminus of mEGFP-ADRB2 locus.</td>
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</tr>
<tr>
<td>mEGFP_N-term_f</td>
<td>ATGGTGAGCAAGGGCGAGGA</td>
<td>Forward primer for generation of partially single-stranded hybrid HDR donor, binds N-terminus of mEGFP</td>
</tr>
<tr>
<td>mEGFP_C-term_linker_r</td>
<td>tctagaCTTGTACAGCTCGTCC</td>
<td>Reverse primer for generation of partially single-stranded hybrid HDR donor, binds C-terminus of mEGFP including a linker sequence</td>
</tr>
<tr>
<td>b2AR-hum_120bp_5'arm_f1</td>
<td>CCCACACCACAGCCGCTGAA</td>
<td>Forward primer for generation of partially single-stranded hybrid HDR donor, binds to 5'UTR of ADRB2</td>
</tr>
<tr>
<td>b2AR-hum_120bp_3'arm_r1</td>
<td>AGACATGACGATGCCCATGC</td>
<td>Reverse primer for generation of partially single-stranded hybrid HDR donor, binds to N-terminus of ADRB2</td>
</tr>
<tr>
<td>b2AR-hum_mEGFP-ADRB2_f1</td>
<td>ATTGGCCGAAAGTTCCCGTA</td>
<td>Forward sequencing primer: verification of endogenous insertion of HDR donor template at ADRB2 locus, binds to 5'UTR of ADRB2 upstream of 5'-homology arm of HDR donor</td>
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<tr>
<td>b2AR-hum_mEGFP-ADRB2_r1</td>
<td>GTCCAAAACCTCGACACAGAA</td>
<td>Reverse sequencing primer: verification of endogenous insertion of HDR donor template at ADRB2 locus, binds to N-terminus of ADRB2 downstream of 3'homology arm of HDR donor</td>
</tr>
<tr>
<td>2nd qPCR_GFP_Fwd</td>
<td>TCGAGAAAGACCCCAACGAG</td>
<td>Forward qPCR primer for amplification of mEGFP-ADRB2. Binds to C-terminus of mEGFP</td>
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<tr>
<td>2nd qPCR_ADRB2_Rev</td>
<td>CGCATGGGCTCTATGTTGGTG</td>
<td>Reverse qPCR primer for amplification of mEGFP-ADRB2. Binds to N-terminus of ADRB2</td>
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**CRISPR strategy**
Efficient endogenous mEGFP-tagging at the N-terminus of ADRB2 in hiPSC was achieved using CRISPR/Cas9 and partially single-stranded, dsDNA-based hybrid homology-directed repair (HDR) donors with symmetric homology arms as previously described [27]. First, a sequence-verified HDR donor plasmid was designed and synthesized (Genscript) including the mEGFP coding sequence flanked by 230 bp homology arms (refer to “Oligonucleotides”). Two different primer pairs were used to generate two PCR products from this donor plasmid encoding for mEGFP alone and for
mEGFP plus 120 bp 5’- as well as 3’-homology arms, respectively (refer to “Oligonucleotides”). A subsequent melt and anneal reaction led to the generation of the partially single-stranded hybrid donor cocktail with HDR donors containing symmetric homology arms and (Supplemental Figure 2A). This cocktail was provided to the cells together with a ribonucleoprotein complex consisting of crRNA, tracrRNA and Cas9 protein (IDT, refer to “Oligonucleotides” for crRNA sequence) via nucleofection using a 4D nucleofector (Lonza). Next, we generated single cell clones using automated hiPSC single cell seeding and clonal expansion using the iotaSciences IsoCell platform [28]. We screened for mEGFP knock-in at the ADRB2 locus by PCR genotyping and selected one homozygous (16) and one heterozygous (66) clone for further experiments.

A 2nd CRISPR round was performed on these two mEGFP-ADRB2 CRISPRed clones 16 and 66 in order to introduce the influenza hemagglutinin signal sequence (HA-SS) (MKTIIALSYIFCLVFA [29]) at the N-terminus of mEGFP-ADRB2. For this, a single-stranded oligodeoxynucleotide (ssODN) encoding for the HA-SS as well as 60bp homology arms was designed and synthesized (IDT) and provided to the cells together with a novel crRNA as mentioned above. Again, single cell clones were generated as described above and screened for homozygous (16-31) and heterozygous (66-35) hiPSC clones via restriction fragment length polymorphism (RFLP) based on the BbsI restriction enzyme cut site introduced by the HA-SS.

Contractility
Cells were imaged in 6-well dishes in brightfield mode on a custom build Zeiss setup. Time series videos of 3-5 min length (with 5 frames per second) were acquired before treatment, after antagonist incubation (100 nM ICI-118551, 300 nM CGP-20712A, 10 µM propranolol) and within 30 min after agonist treatment (1 µM isoproterenol). Contraction rates and other parameters were extracted from videos using the ImageJ software “Myocyter v1.3” [30].

qRT-PCR
Cells were lysed and RNA extracted according to the RNeasy Mini kit (Qiagen). The cDNA synthesis was performed using the SuperScript III First-Strand Synthesis SuperMix (Invitrogen). Finally, qRT-PCR was done using the QuantiTect SYBR Green PCR kit (Qiagen) on an QuantStudio 6 flex qPCR device (Thermo Fisher Scientific). Primers for GAPDH (Geneglobe ID: QT00079247) and ADRB2 (Geneglobe ID: QT00200011) were purchased from Qiagen and primers for mEGFP-ADRB2 were self-made (refer to “Oligonucleotides”).

Cell Preparation for Confocal Imaging
Three days prior to imaging hiPSC-CM were detached as single cells and seeded in Geltrex-coated, eight-well, glass-bottom µ-slides (Ibidi). Until the day of experiment hiPSC-CM were cultured in maintenance medium with 3 µM CHIR99021 and 1x RevitaCell supplement (Life Technologies). H9c2 cells were seeded in eight-well glass-
bottom μ-slides (Ibidi) without coating and transfected 24 h later using Lipofectamine 2000 (Thermo Fisher Scientific) according to the manufacturer’s instructions. Two days post transfection H9c2 cells and also hiPSC-CM were imaged in imaging buffer (pH 7.4, 20 mM 4- (2-hydroxyethyl)-1-piperazineethanesulfonic acid [Hepes], 137 mM NaCl, 5 mM KCl, 1 mM MgCl₂, 1 mM CaCl₂, 0.5% bovine serum albumin [BSA]). Prior to imaging, hiPSC-CMs were preincubated for 40 min to 1 h with either 100 nM CGP-20712 (to image β₂-AR) or 50 nM ICI-118,551 (to image β₁-AR) diluted in the imaging buffer. Then, 50 nM JE1319 ligand was diluted in imaging buffer and directly added to the cells (with CGP-20712 or ICI-118,551 as described) for 40 min to 1 h. During the last 15 min of ligand incubation, 0.25x CellMask Green Plasma Membrane Stain (Invitrogen) was added to the hiPSC-CMs. After all incubations, hiPSC-CMs were washed three times using imaging buffer and were imaged in imaging buffer containing the respective β-AR antagonist as well as 50 μM para-aminoblebbistatin (Optopharma) to inhibit spontaneous contractions. Finally, cells were imaged on a Leica SP8 WLL confocal laser scanning microscope under physiological conditions (37 °C, 5% CO₂, 85% humidity) using a sample incubator (Stage Top Chamber; OKOlab).

Immunofluorescence
At the day of fixation hiPSC-CM were washed with 250 µl phosphate buffered saline (PBS) (PAN-Biotech) per 8-well μ-slide (Ibidi). Cells were fixed using 4% paraformaldehyde for 30 min at room temperature (RT). Afterwards cells were washed again with PBS and stored at 4°C. Cells were blocked using NDS buffer (10% NDS, 0.2% BSA, 0.3% TritonX100) for 1 h at RT and directly labelled overnight at 4°C with primary antibodies (1:400 mouse mAB for sarcomeric alpha-actinin [CloneEA-53, Sigma-Aldrich, A7811] and 1:200 rabbit pAB for cardiac Troponin T [Abcam, ab45932]). After washing cells once with PBS, secondary antibodies were applied for 2 h in the dark (1:250 anti rabbit IgG coupled to AlexaFluor-555 [Invitrogen, A31572] and 1:250 anti mouse IgG coupled to AlexaFluor-488 [Invitrogen, A21202]). Two drops of NucBlue™ Fixed Cell ReadyProbes™ (Invitrogen, R37606) were added to the secondary Ab solution to stain for nuclei. Cells were washed twice with PBS and then imaged on a Leica SP8 WLL.

Acquisition of line scans
In order to extract the behavior of the diffusing species linescan-FCS (Fluorescence Correlation Spectroscopy) was performed in which a confocal beam is repeatedly scanned at high speed over the same portion of the sample. The resultant recorded kymographs or linescans are further analyzed by a custom-implemented code in MATLAB to extract information about diffusion data of receptors from autocorrelation functions (as previously described in [15]).

In short, linescans were acquired on a commercial confocal laser scanning microscope, Leica SP8 WLL, with a white light laser (WLL). Via the HC PLAP CS2 40x1.3 NA oil immersion objective (Leica) a pixel size of 50 nm was used to acquire about 6 x 10⁵ lines of 256 pixels each at a speed of 1800 Hz. In order to stabilize the focal position,
the IR laser based autofocus of the microscope (Leica, Adaptive Focus Control) was enabled while the ligand JE1319 (Alexafluor 647-based) was excited at a wavelength of 633 nm with 5% or 50% laser power which upon calibration corresponded to total power outputs in the µW-range. Hybrid detectors (HyD) in photon counting mode were applied to detect emission in the range of 650-751 nm. The beam waist was extracted by observation of the profiles of fluorescent microspheres (Tetraspeck, Thermo Fisher Scientific) as in [15] resulting in a lateral waist of $\omega_0^{(633 \text{ nm})} = 0.33 \, \mu\text{m}$.

Analysis of confocal linescans
Confocal linescans were analyzed as previously described [13, 15]. Briefly, fluorescence correlation spectroscopy (FCS) extracts information on equilibrium processes in the sample by comparing statistical fluctuations of fluorescent particles in an effective detection volume. Linescan FCS greatly increases the statistical accuracy of these measurements due to the repeated scanning of one line. The recorded time-trace of fluorescent intensity (I) fluctuations in the effective detection volume allows for the calculation of an autocorrelation $G(\tau)$ reflecting the time-scale of these fluctuations.

$$G(\tau) = \frac{\langle I(t) \cdot I(t + \tau) \rangle}{\langle I(t) \rangle^2}$$

The pointed brackets represent the average over all time values t. In order to extract diffusion parameters, such as the diffusion time $\tau_D$ or the diffusion constant D (which are connected via $\tau_D = \frac{\omega_0^2}{4D}$), the autocorrelation function $G(\tau)$ is fit to the two-dimensional model of the autocorrelation function.

$$G_{2D}(\tau) = \frac{1}{N} \frac{1}{1 + \frac{4D\tau}{\omega_0^2}} + G_\infty$$

Where $\tau$ represents the time lag, N the average number of particles and $G_\infty$ the limiting value of $G(\tau)$ for $\tau \to \infty$. $\omega_0$ is the beam waist, and describes the extent of the effective detection volume in the focal plane where the intensity has dropped to $e^{-2}$.

In particular, an initial removal of the first 180225 lines of each scan reduces the influence of photo bleaching. The analysis is further corrected for bleaching via a moving average. A high pass filter reduces slow fluctuations by dividing the total time into smaller pieces of about 36:25 s.

Each autocorrelation curve is normalized to the curve’s $G(0)$. In the case that no correlation was extracted the curve is normalized to its highest peak.
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Author contributions

P.G. and P.A. conceived the study and designed research. P.G. performed research with support from N.T. on generation of CRISPR clones and stem cell handling/differentiation. P.G. analyzed data. M.B.-P. analyzed linescan-FCS data and provided the software for this. P.G. and M.B.-P. prepared the figures. P.A. drafted the manuscript with contributions from P.G. and M.B.-P. The manuscript was edited by P.G., M.B.-P. and N.T. All authors reviewed and approved the final manuscript. P.A. initiated and supervised the research. P.A. and M.J.L. acquired funding.
Supplemental Figure 1 Timeline displaying differentiation and maturation of hiPSC-CMs. (A) Shows changes of cellular morphology of undifferentiated WT hiPSC (day 0) and WT cells differentiated and matured for 7, 21, 57 and 94 days in culture. Scale bar is 30 µm. (B) Differentiation strategy based on small molecules (see Methods) depicting the most important treatment media indicated by day and duration of treatment and days of sample collection (25 +/-4 days; 60 +/-4 days; 100 +/-7 days).
Supplemental Figure 2 Strategy for generation and identification of the endogenously CRISPRed HA-mEGFP-ADRB2 clones. (A) Schematic of the 2-step CRISPR approach for N-terminal mEGFP-tagging of ADRB2 based on partially single-stranded hybrid donors with symmetric homology arms for the 1st homology-directed repair (HDR). In a 2nd CRISPR step the influenza hemagglutinin signal sequence (HA-SS) (MKTIALSYIFCLVFA [29]) was introduced endogenously at the N-terminus of mEGFP-ADRB2 for improved membrane targeting [31] using a single-stranded oligodeoxynucleotide (ssODN) as HDR donor. (B) PCR genotyping of 16 CRISPRed isogenic hiPSC clones on agarose gel after 1st homology-directed repair based on genomic PCR using primers that bind outside the homology arms of the HDR donor as indicated with blue arrows in (A). Homozygous (clone 16) and heterozygous (clone 66) clones are highlighted in the yellow rectangle. (C) Genotyping of two isogenic HA-mEGFP-ADRB2 CRISPRed clones 16-31 and 66-35 which were derived from mEGFP-ADRB2 CRISPRed clones 16 and clone 66 (highlighted in (B)) after using them for the 2nd CRISPR step (introducing HA-SS). Clones were identified by restriction fragment length polymorphism (RFLP) using the BbsI restriction enzyme cut site in the HA-SS (highlighted in orange in A). Shown are undigested (undig) and BbsI-digested (dig) genomic PCR reactions and the expected resulting PCR fragment sizes. (D) Shows DNA chromatograms of clones 16-31 and 66-35 based on Sanger sequencing at their CRISPRed ADRB2 loci verifying correct in frame integration of the HA-mEGFP sequence at the N-terminus of ADRB2.
Supplemental Figure 3 (A) Mean diffusion constants and (B) mean number of particles extracted from the autocorrelation curves in Figure 3. Error bars represent SD values. Dots and small squares indicate diffusion constants and receptor numbers extracted at day 107 (and are not part of the day 100 mean).

Supplemental Figure 4 HA-mEGFP-ADRB2 overexpression in H9c2 cells. (A) Confocal micrograph of the basolateral membrane of an H9c2 cell expressing the construct HA-mEGFP-ADRB2, excited at 488 nm. (B) corresponding DIC image. Scale bar is 20 µm.

Video S1 Spontaneous contractions of the hiPSC-CM clone 16-31 (at day 100) before β2-AR stimulation and after 25 min of 300 nM CGP-20712A addition

Video S2 Spontaneous contractions of the hiPSC-CM clone 16-31 (at day 100) after 5 min of β2-AR stimulation
References