Product description	Human iPSC clonal line in which MYL7 has been endogenously tagged with mEGFP using CRISPR/Cas9 technology	
Parental cell line	Parental hiPSC line (WTC/AICS-0 at passage 33) derived from fibroblasts reprogrammed using episomal vectors (OCT3/4, shp53, SOX2, KLF4, LMYC, and LIN28). Coriell catalog: GM25256	
Publication(s) describing iPSC establishment	Kreitzer et al (2013) Am. J. Stem Cells, 30; 2(2): 119-31	
Passage of gene edited iPSC reported at submission	p39 ^a	
Number of passages at Coriell	0	
Media	mTeSR1	
Feeder or matrix substrate	matrix substrate Matrigel	
Passage method	Accutase	
Thaw	1 million cells (ea vial) in 10 cm plate - ready for passaging in 3-4 days	
Seeding density	ding density 400K cells/10-cm plate every 4 days or 800K cells/10-cm plate every 3 days (sculture protocol)	

Test Description ^b	Method	Specification	Result
Post-Thaw Viable Cell Recovery	hiPSC culture on Matrigel	> 50% confluency 3-4 days post-thaw (10 cm plate)	Pass
mEGFP insertion at genomic locus - precise editing	PCR and Sanger sequencing of recombinant and wildtype alleles	C-term insertion of mEGFP in frame with exact predicted recombinant allele junctions. No additional mutations in either allele.	Pass
Copy number	ddPCR ^c assay for mEGFP and RPP30 reference gene ^d	mEGFP/RPP30: $\sim 0.5 = \text{Mono-allelic}$ $\sim 1.0 = \text{Bi-allelic}$	Mono-allelic (0.48)
Plasmid integration	ddPCR assay to detect plasmid integration into the genome	$\begin{array}{l} {\rm AmpR/RPP30:} \\ < 0.1 = {\rm no~plasmid} \\ {\rm integration} \end{array}$	Pass (0.00)
Mutational analysis	Whole exome sequencing ^f	Check for acquired mutations (not detected in p8 ^a parental line) that: 1) Correspond to off-target sites predicted by Cas-OFFinder ^e 2) Affect genes in Cosmic Cancer Gene Census	Sequencing planned
mEGFP localization	Spinning Disk confocal live cell imaging	Localization to sarcomeres in hiPSC-derived cardiomyocytes	Localizes to striations in myofibrils, consistent with localization to thick filaments (myosin-based) within the sarcomere and exclusion from the Z-disk and I-band.
Expression of tagged protein	Western blot	Expression of expected size product	Not performed

Growth rate	ATP quantitation ^g	Comparable to parental line	Pass (measured at p39) ^a
Expression of stem cell markers	Flow cytometry	Transcription factors: $ \begin{array}{l} \text{OCT4/SOX2/NANOG} \geq \\ 85\% \\ \text{Surface markers:} \\ \text{SSEA3, TRA-1-60} \geq 85\%; \\ \text{SSEA1} \leq 15\% \\ \end{array} $	Pass
Germ layer differentiation	Trilineage differentiation ^h as assayed by ddPCR gene expression analysis	Expression of endoderm (SOX17), mesoderm (Brachyury), and ectoderm (PAX6) markers upon directed differentiation to all three germ layers	Pass
Cardiomyocyte differentiation	Modified small molecule differentiation (Lian et al. 2012) ⁱ	Beating initiated (D7-D14) and Cardiac Troponin T expression (D12-D30) by flow cytometry	Pass
Karyotype	G-banding (30 cell analysis)	Normal karyotype, 46 XY	Pass
Mycoplasma	qPCR (IDEXX)	Negative	Pass
Sterility (bacterial, yeast and fungal testing)	Direct inoculation and incubation for 10 days	No growth after 10 days	Pass
Viral Panel Testing ^j	PCR	Negative when assayed for CMV, EBV, HepB, HepC, HIV1, and HPV	Pass
$\begin{array}{c} \textbf{Identity of} \\ \textbf{unedited parental} \\ \textbf{line}^k \end{array}$	STR	29 allelic polymorphisms across 15 STR loci compared to donor fibroblasts	Identity matched

^a This is the number of passages beyond the original parental line (WTC/AICS-0 at passage 33).

^b All QC assays are performed on stem cells except when noted otherwise.

^c Droplet digital PCR using Bio-Rad QX200

^d RPP30 is a reference 2 copy gene used for normalization.

 $^{^{\}mathrm{e}}$ Bae et al (2014) Bioinformatics. 30(10): 1473-1475

^f Nextera rapid capture exome

 $^{^{\}rm g}$ Promega Cell
Titer-Glo Luminescent Cell Viability Assay (Catalog #G7571)

 $^{^{\}rm h}$ STEMCELL Technologies STEMdiff Trilineage Differentiation Kit (Catalog #05230)

ⁱ Lian et al (2012) PNAS. 109(27):E1848-E1857

^j Viral panel testing was conducted for the parental WTC line prior to editing. Sterility (bacterial, fungal) and mycoplasma testing were conducted in both the parental and edited lines.

^k STR tests were conducted for the WTC parental line prior to editing. WTC is the only cell line used by AICS. Edited WTC cells were not re-tested because they did not come into contact with any other cell lines.

mEGFP tagging strategy: Used CRISPR-Cas9 methodology to introduce mEGFP at C-terminus of MYL7 as shown below.

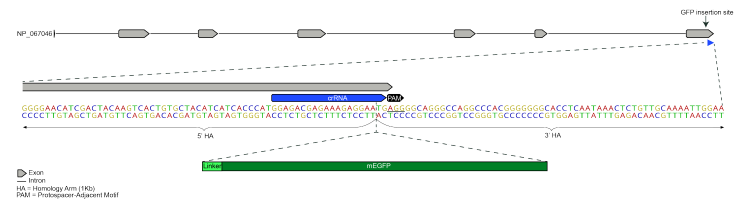


Figure 1: Top: MYL7 locus; Bottom: Zoom in on mEGFP insertion site at MYL7 C-terminus; For more information on tagging transcriptionally silent genes in hiPSCs see: Roberts et al. 2018. Scarless gene tagging of transcriptionally silent genes in hiPSCs to visualize cardiomyocyte sarcomeres in live cells. bioRxiv doi:10.1101/342881

Post-thaw imaging: One vial of distribution lot was thawed (cells were treated with ROCK inhibitor for 24hrs post-thaw refer to culture protocol). Cultures were observed daily. Colonies were photographed one and three days post-thaw^{1,2} using a Leica microscope at 4X and 10x magnification.

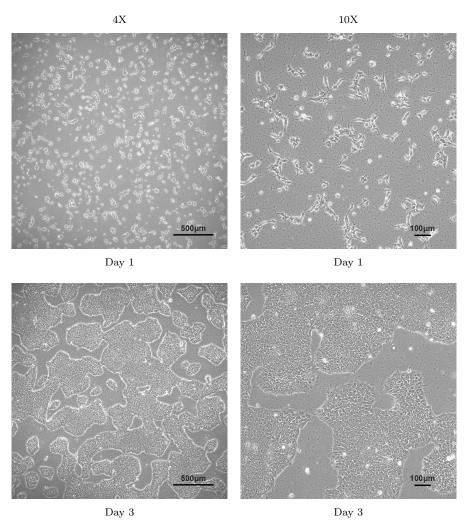


Figure 2: Viability and colony formation one day and three days post-thaw

 $^{^1\}mathrm{Cells}$ may take up to 3 passages to recover after thaw

 $^{^2\}mathrm{Morphologies}$ observed post-thaw are representative of cell morphologies observed post-passage

Imaging labeled structures in endogenously tagged hiPSC derived cardiomyocytes: The tagged proteins are expressed endogenously and therefore may not appear as bright as they would in an overexpressed system. For imaging, we plate cells onto high-quality glass bottom 24-well plates (Cellvis) coated with 0.1% w/vol polyethylenimine (PEI) and $25\mu g/ml$ laminin. Cells are imaged in phenol red-free RPMI 1640 media (Gibco) supplemented with B-27 containing insulin (Gibco). Our most common microscope configuration is a Zeiss spinning disk fluorescence microscope with a Yokogawa CSUX1 head, Hamamatsu CMOS camera, and a 488 laser (GFP). Cells are imaged with a 40x 1.2 NA water immersion objective at 37°C and 5% CO₂ in a temperature-controlled chamber. The approximate laser power measured at the sample for our standard 100x images is \sim 2.5 mW.

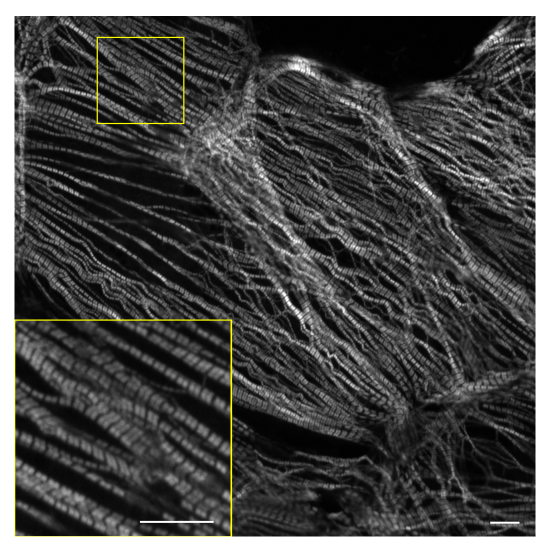


Figure 3: Single, mid-level plane of live hiPSC-derived cardiomyocytes expressing mEGFP-tagged MLC-2a protein. Twelve days after the onset of differentiation, cells were plated on PEI and laminin coated glass and imaged in 3D on a spinning disk confocal microscope 20 days later (32 days total after the onset of differentiation). Scale bars, $10 \mu m$.