Microgravity-induced stress mechanisms in human stem cell-derived cardiomyocytes

Aviseka Acharya, Harshal Nemade, Symeon Papadopoulos, ... Athanasia Mizi, Argyris Papantonis, Agapios Sachinidis

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Highlights

- Simulated microgravity (SMG) causes ROS production in human cardiomyocytes (CMs)
- SMG inhibits mitochondria function and energy metabolism and induces senescence of CMs
- SMG attenuates contractile velocity, beating frequency and Ca²⁺ influx in CMs
- SMG induces chromosomal changes and modifies the chromosomal architecture in CMs
Microgravity-induced stress mechanisms in human stem cell-derived cardiomyocytes

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SUMMARY

Exposure to outer space microgravity poses a risk for the development of various pathologies including cardiovascular disease. To study this, we derived cardiomyocytes (CMs) from human-induced pluripotent stem cells and exposed them to simulated microgravity (SMG). We combined different “omics” and chromosome conformation capture technologies with live-cell imaging of various transgenic lines to discover that SMG impacts on the contractile velocity and function of CMs via the induction of senescence processes. This is linked to SMG-induced changes of reactive oxygen species (ROS) generation and energy metabolism by mitochondria. Taken together, we uncover a microgravity-controlled axis causing contractile dysfunctions to CMs. Our findings can contribute to the design of preventive and therapeutic strategies against senescence-associated disease.

INTRODUCTION

Gravity on Earth has had a documented impact on the development of organisms and is essential for sustaining human life (Demontis et al., 2017; Morey-Holton, 2003). However, life outside the Earth has always intrigued the human mind. Recently, the deep space exploration of Mars have significantly enhanced research on the influence of microgravity on the human body. Over the last 50 years, gravitational biology-related investigations have revealed significant effects of microgravity on organ development using animal models (Ruden et al., 2018). Currently, international space agencies and private corporations are planning missions to the Moon and Mars that require the long presence of humans in space (for a review see ref. (Hughson et al., 2018)).

Studies on astronauts who had remained in space for prolonged periods of time revealed two main risk factors for developing various diseases, including cardiovascular diseases: microgravity and space radiation. In particular, such studies proved that drastic reduction below Earth gravity levels (1 g) leads to significant health problems to humans due to their adaptation to outer space microgravity environments (Hughson et al., 2018; Patel, 2020), acting as a major stress stimulus for astronauts. In this context, astronauts are exposed to microgravity for several weeks face such challenges to their health as loss of muscle and bone mass (Demontis et al., 2017; Pelligr et al., 2020), impaired immune defense (Akiyama et al., 2020; Crucian et al., 2018), hematic, metabolic and endocrine complications (Demontis et al., 2017; Pelligr et al., 2020), brain and behavioral complications (Hupfeld et al., 2021), as well as cardiovascular problems (Hughson et al., 2018; Patel, 2020). It therefore becomes obvious that microgravity-related complications are reminiscent of cellular aging processes, in particular of cellular senescence due to reactive oxygen species production (Demontis et al., 2017; Strollo et al., 2018; Vernikos and Schneider, 2010).

Space medicine research, specifically in respect to cardiovascular studies, is of fundamental importance for developing health care interventions for humans with a prolonged exposure to microgravity (Afshinnekoo et al., 2020). To this end, understanding microgravity-induced health complications is a precondition for developing successful therapeutic strategies for humans. Several methodologies have been established for performing in vivo (Pandiarajan and Hargens, 2020; Watenpaugh, 2016) and in vitro investigations (Kiss et al., 2019) on Earth under artificial microgravity conditions. Such microgravity investigations can be performed using traditional real and simulated microgravity platforms such as drop towers, parabolic flights, sounding rockets and clinostats (Amselem, 2019). However, except for the clinostat platform that allows exposure of cell systems for longer times (up to several days), all other methods induce microgravity acutely for a few seconds to minutes.
Due to ethical concerns (Sachinidis et al., 2019), an additional obstacle in ascertaining microgravity mechanisms that are of relevance for humans has been the availability and access to “normal” human cellular systems. Nowadays, human-induced pluripotent stem cell (hiPSC)-derived somatic cells alleviate this obstacle and offer an unlimited source of functional cells, including synchronously beating cardiomyocytes, possessing the key functional properties of cardiomyocytes isolated from human hearts.

To date, because of several experimental limitations, in particular the very short microgravity time exposure of 20 s, deeper insights into the molecular mechanisms induced by microgravity and promoting heart diseases were not possible (Acharya et al., 2019). Here, we address this shortcoming by describing the molecular mechanisms and key features of microgravity-induced senescence in human iPSC-derived cardiomyocytes (CMs) by a combination of high-throughput and functional studies, made possible by their controlled exposure to simulated long-term microgravity (SMG) under 1 g laboratory conditions.

RESULTS
Gene expression changes in CMs exposed to SMG at the transcriptome and proteome levels
Purified CMs isolated from hiPSC were exposed to SMG for 48 h (hereafter referred to as “SMG-CMs”). SMG was induced using a slide-flask 2D clinostat device (see STAR Methods for details, Figure S1). Control cardiomyocytes are hereafter referred to as “c-CMs”. To assess potential changes in gene expression in CMs exposed to SMG for 48 h, we isolated total cell RNA from SMG- and c-CMs and subjected it to high-throughput sequencing RNA sequencing (RNA-seq). Following data analysis, 825 genes were statistically differentially expressed (DEGs) due to SMG exposure. Of these, 68% (n = 562) were significantly upregulated and 263 downregulated (Table S1). A query of gene ontology (GO) terms and KEGG/Reactome pathways related to these DEGs via Meta- scape (Zhou et al., 2019) suggested that SMG promotes adhesion of CMs by upregulating extracellular matrix (ECM) genes (top terms listed in Table 1; complete GO term lists for up- and downregulated genes are shown in Tables S2 and S3, respectively). Moreover, SMG induced hypoxia as cardiomyocytes-expressed genes countering the drop of intracellular oxygen levels. Accordingly, SMG-CMs increased their glycolytic metabolism (as evidenced by induced pyruvate metabolic processes), as well as their contractile activity via genes involved in Ca2+ homeostasis. On the other hand, the 263 significantly downregulated genes in SMG-CMs could be linked to reduced ATP production via the inhibition of oxidative phosphorylation. Together, these data suggest that SMG exposure for just 48 h suffices for inducing cellular stress to CMs, primarily reflected in changes to the transcriptome that pertain to mitochondrial structure and functions.

Next, we applied quantitative SILAC-based mass spectrometry to SMG- and c-CMs to capture proteins differentially expressed by SMG. We identified 56 down- and 123 upregulated proteins upon 48 h of microgravity exposure (listed in Table S4). Metascape analysis of GO terms/pathways linked to SMG-upregulated proteins showed that DNA packaging (mostly via histone level changes), chromatin organization, and epigenetic modifiers are strongly affected (Table 2). Notably, SMG also promoted protein expression relevant for oxidative stress-induced senescence, in line with the mito-centric transcriptional changes recorded by RNA-seq (above), as well as changes to ECM components. Proteins downregulated as a result of SMG exposure could be linked to the control of CM growth and to mitophagy (complete GO term lists for up-/downregulated proteins are shown in Tables S5 and S6, respectively). Thus, this proteome-wide analysis also highlights SMG-induced stress to CMs.

Spatial chromosome reorganization follows CM exposure to SMG
Combined, the transcriptome and proteome changes we recorded hinted toward changes in chromosomal conformation. We therefore investigated 3D chromatin architecture changes in SMG-CMs and c-CMs using chromosome conformation capture technology under native conditions (iHi-C). To this end, and given the limited availability of CMs in our SMG experimental set up, we opted for our modified iHi-C 2.0 approach (Mizi et al., 2020) that omits cell fixation, but also incorporates the “easy Hi-C” workflow tailored to low input samples (Lu et al., 2020). Following a novel way for Hi-C data analysis (see STAR Methods for details) and the comparison of the SMG- and c-CM iHi-C matrices, we could identify changes across all levels of 3D genome organization.

First, looking into the organization of the transcriptionally active A- and mostly inactive B-compartments, we found that ~12% switched from A to B, and 6% from B to A. However, these multi-Mbp domains only hosted few of the DEGs recorded above (Figure 1A). Second, we investigated Mbp-sized topologically associating domains (TADs), commonly viewed as the “building blocks of chromosomes” (Beagan and Phillips-Cremins, 2020). Here, >35% of TADs in c-CMs are remodeled in SMG-CMs, with the majority (~25%)
Table 1. Metascape analysis of the upregulated and downregulated genes in SMG-CMs versus the c-CMs; representative BPs, KEGG pathways, and Reactome Gene Sets

### Upregulated: Representative enriched BPs and pathways stimulated by SMG

| Term                      | Description                                      | LogP  | Log(q-value) | InTerm_InList |
|---------------------------|--------------------------------------------------|-------|--------------|---------------|
| GO:0030198                | extracellular matrix organization                | -32,4 | -28,6        | 54/395        |
| GO:0070482                | response to oxygen levels                        | 26,0  | -22,4        | 47/385        |
| GO:0001666                | response to hypoxia                              | -24,0 | -20,6        | 43/348        |
| GO:0036293                | response to decreased oxygen levels              | -23,4 | -20,0        | 43/360        |
| GO:2000147                | positive regulation of cell motility             | -15,6 | -12,8        | 43/579        |
| GO:006090                 | pyruvate metabolic process                       | -15,0 | -12,2        | 23/153        |
| GO:007307                 | heart development                                | -14,7 | -12,0        | 41/556        |
| GO:006096                 | glycolytic process                               | -14,4 | -11,7        | 20/114        |
| GO:031589                 | cell-substrate adhesion                          | -13,8 | -11,2        | 32/360        |
| GO:006178                 | glucose catabolic process to pyruvate            | -12,6 | -10,1        | 11/27         |
| GO:000160                 | cell-matrix adhesion                             | -7,9  | -5,8         | 19/231        |
| GO:0006874                | cellular calcium ion homeostasis                 | -6,9  | -4,9         | 25/443        |
| GO:0055074                | calcium ion homeostasis                          | -6,7  | -4,7         | 25/455        |
| GO:0060047                | heart contraction                                | -6,6  | -4,7         | 19/277        |

**KEGG Pathway**

| Term                      | KEGG Pathway | LogP  | Log(q-value) | InTerm_InList |
|---------------------------|--------------|-------|--------------|---------------|
| ko04512                   | ECM-receptor interaction                          | -18,7 | -15,6        | 21/82         |
| ko04510                   | Focal adhesion                                     | -17,3 | -14,4        | 28/199        |
| ko04151                   | PI3K-Akt signaling pathway                         | -12,1 | -9,6         | 29/342        |
| hsa04066                  | HIF-1 signaling pathway                            | -11,9 | -9,5         | 18/119        |
| ko000110                  | Glycolysis/Gluconeogenesis                         | -10,2 | -7,9         | 13/67         |

**Reactome Gene Sets**

| Term                      | R-HSA-1474244 | Extracellular matrix organization | LogP  | Log(q-value) | InTerm_InList |
|---------------------------|---------------|----------------------------------|-------|--------------|---------------|
| R-HSA-1474290             | Collagen formation | -33,1                           | -29,1 | 49/301       |
| R-HSA-216083              | Integrin cell surface interactions                | -19,7 | -16,5        | 22/85         |
| R-HSA-3000178             | ECM proteoglycans                                   | -19,5 | -16,3        | 21/76         |
| R-HSA-1474228             | Degradation of the extracellular matrix            | -18,1 | -15,0        | 25/140        |
| R-HSA-8948216             | Collagen chain trimerization                       | -17,2 | -14,3        | 16/44         |

### Downregulated (representative enriched BPs and pathways suppressed by SMG)

| Term                      | Description                                      | LogP  | Log(q-value) | InTerm_InList |
|---------------------------|--------------------------------------------------|-------|--------------|---------------|
| GO:0006119                | oxidative phosphorylation                         | -33,9 | -30,0        | 32/148        |
| GO:0046034                | ATP metabolic process                             | -26,7 | -23,6        | 35/313        |
| GO:0042773                | ATP synthesis coupled electron transport          | -26,7 | -23,5        | 24/100        |
| GO:0042775                | mitochondrial ATP synthesis coupled electron transport | -26,8 | -23,6    | 24/99         |
| GO:0007005                | mitochondrial organization                         | -15,0 | -12,5        | 31/551        |
| GO:0006123                | mitochondrial electron transport, cytochrome c to oxygen | -12,9 | -10,6        | 9/21          |
| GO:0019646                | aerobic electron transport chain                   | -12,9 | -10,6        | 9/21          |
| GO:1902600                | proton transmembrane transport                    | -12,9 | -10,5        | 17/157        |
| GO:0006120                | mitochondrial electron transport, NADH to ubiquinone | -10,1 | -7,8        | 10/55         |
| GO:0010257                | NADH dehydrogenase complex assembly               | -9,4  | -7,1         | 10/65         |
| GO:004207                 | cristae formation                                 | -6,1  | -4,0         | 6/35          |

**KEGG Pathway**

| Term                      | KEGG Pathway | LogP  | Log(q-value) | InTerm_InList |
|---------------------------|--------------|-------|--------------|---------------|
| hsa000190                 | Oxidative phosphorylation                          | -29,2 | -25,9        | 30/167        |

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splitting into smaller TADs, with ∼16% merged into larger TADs (Figure 1B). Third, by plotting the decay of the interaction frequency of chromatin fragments as a function of genomic distance, we observed that longer-range interactions (>20 Mbp in separation) are strengthened upon SMG exposure, potentially indicative of widespread chromosome compaction. At the same time, more proximal interactions are markedly weakened in response to SMG (Figures 1C and 1D). Finally, in an attempt to link 3D chromatin reorganization with SMG-driven gene expression changes, we plotted mean iHi-C signal changes in the 1 Mbp around up-/downregulated gene TSSs. For upregulated TSSs, we observed the focal gain of encompassing interactions, potentially indicative of inclusion in novel sub-Mbp (i.e., TAD-sized) domains (Figure 1E, left panel). For downregulated TSSs, there was a marked gain in shorter (<250-kbp) contacts, again potentially indicative of local chromatin compaction (Figure 1E, right panel).

**SMG impacts on the sarcomeric and cytoskeletal structure alignment in CMs**

The transcriptome and SILAC-proteome data suggested that SMG modifies CM contractility as well as modify their ECM. Based on these observations, we investigated the effects of SMG on sarcomere assembly by immunostaining for cardiac α-Actinin, as well as by transmission electron microscopy (TEM). Although no significant changes in the morphology of SMG-CM sarcomeres were observed (Figures 2A), TEM revealed the formation of stress fibers and membrane caveolae in a subset of SMG-CMs (Figure 2B). Such fibers and caveolae are characteristic changes to the cell membrane due to mechanical stress (Echarri and Del Pozo, 2015). The formation of caveolae in SMG-CMs could be also confirmed by Western blotting showing higher levels of CAV1 as compared to c-CMs (Figure 2C). These findings suggest that although SMG exposure does not lead to significant sarcomere disassembly, it does induce stress fiber and caveolae in SMG-CMs.

**Effects of SMG on mitochondrial structure, function, and ATP synthesis**

Given the physiology of CMs, focused on executing synchronous beating, the formation of caveolae and stress fibers is possibly indicative of effects on contractility. To assess this, we further evaluated different aspects of mitochondrial function under SMG (as also hinted by our Metascape analyses; see Tables 1 and 2). First, in order to monitor ROS in the mitochondria, we generated DyRed1-E5⁺-CMs after differentiation of DyRed1-E5⁺-hiPSCs and exposed them to SMG for 48 h. DyRed1-E5 is a mutant form of the DsRed protein that is mitochondrial-targeted and sensitive to free radicals (Figure S2). It changes its fluorescence from green to red in response to increased ROS levels and/or to high mitochondrial turnover. In positive control experiments, DyRed1-E5⁺ c-CMs were treated with H₂O₂, which increases intracellular ROS. This resulted in almost complete elimination of green fluorescent signal from the cells (Figure 3B). Similarly, high-resolution image analysis of DyRed1-E5⁺ SMG-CMs also revealed mitochondria with intense red fluorescence indicative of accumulating of ROS in response to SMG (see representative Video S1), which was not the case for DyRed1-E5⁺ c-CMs (see representative Video S2 and Figure 3B).

| Term                     | Description                                                                 | LogP | Log(q-value) | InTerm_InList |
|--------------------------|------------------------------------------------------------------------------|------|--------------|---------------|
| hsa03010                 | Ribosome                                                                     | −21,3| −18,5        | 24/163        |
| hsa04260                 | Cardiac muscle contraction                                                  | −15,6| −13,1        | 16/91         |
| hsa_M00154               | Cytochrome c oxidase                                                         | −12,3| −10,0        | 9/24          |
| R-HSA-163200             | Respiratory electron transport, ATP synthesis by chemiosmotic coupling, and heat production by uncoupling proteins | −33,3| −29,5        | 30/125        |
| R-HSA-1428517            | The citric acid (TCA) cycle and respiratory electron transport               | −28,5| −25,2        | 30/176        |
| R-HSA-2262752            | Cellular responses to stress                                                 | −21,4| −18,6        | 39/584        |
| R-HSA-3700989            | Transcriptional Regulation by TP53                                            | −16,0| −13,5        | 27/365        |
| R-HSA-5628897            | TP53 Regulates Metabolic Genes                                               | −14,5| −12,1        | 15/87         |
| R-HSA-69278              | Cell Cycle, Mitotic                                                          | −11,5| −9,2         | 27/561        |
| R-HSA-8949613            | Cristae formation                                                            | −6,4 | −4,3         | 6/31          |
Table 2. Metascape analysis of the SILAC-upregulated and downregulated proteins in SMG-CMs versus the c-CMs; Representative BPs, KEGG pathways, and Reactome Gene Sets

**Upregulated:** Representative enriched BPs and pathways stimulated by SMG

| Term                  | Description                                             | LogP | Log(q-value) | InTerm_InList |
|-----------------------|---------------------------------------------------------|------|--------------|---------------|
| GO:0071103            | DNA conformation change                                 | −26.1| −22.8        | 26/323        |
| GO:0071824            | protein-DNA complex subunit organization                | −23.1| −20.3        | 23/282        |
| GO:0006342            | chromatin silencing                                     | −22.2| −19.4        | 18/133        |
| GO:0065004            | protein-DNA complex assembly                            | −21.7| −19.0        | 21/242        |
| GO:0006323            | DNA packaging                                           | −21.4| −18.8        | 20/211        |
| GO:0045814            | negative regulation of gene expression, epigenetic      | −21.4| −18.8        | 18/146        |
| GO:0006333            | chromatin assembly or disassembly                       | −19.2| −16.7        | 18/192        |
| GO:0006281            | DNA repair                                              | −11.0| −8.6         | 18/567        |
| GO:0030198            | extracellular matrix organization                       | −9.2 | −6.9         | 14/395        |
| GO:0042254            | ribosome biogenesis                                     | −7.4 | −5.2         | 11/307        |

**KEGG Pathway**

| Term                  | Description                                             | LogP | Log(q-value) | InTerm_InList |
|-----------------------|---------------------------------------------------------|------|--------------|---------------|
| hsa04217              | Necroptosis                                             | −18.7| −16.2        | 17/168        |
| ko03030               | DNA replication                                         | −10.0| −7.7         | 7/36          |
| ko04512               | ECM-receptor interaction                                | −8.9 | −6.6         | 8/82          |
| hsa04510              | Focal adhesion                                          | −5.6 | −3.5         | 8/215         |
| ko04151               | PI3K-Akt signaling pathway                              | −4.2 | −2.1         | 8/342         |

**Reactome Gene Sets**

| Term                  | Description                                             | LogP | Log(q-value) | InTerm_InList |
|-----------------------|---------------------------------------------------------|------|--------------|---------------|
| R-HSA-3214815         | HDACs deacetylase histones                              | −28.8| −25.2        | 20/94         |
| R-HSA-5334118         | DNA methylation                                         | −26.2| −22.9        | 17/65         |
| R-HSA-212300          | PRC2 methylates histones and DNA                        | −25.3| −22.1        | 17/73         |
| R-HSA-2559586         | DNA Damage/Telomere Stress Induced Senescence           | −24.5| −21.4        | 17/80         |
| R-HSA-4839726         | Chromatin organization                                  | −23.4| −20.5        | 23/274        |
| R-HSA-3214847         | HATs acetylase histones                                  | −23.3| −20.4        | 19/142        |
| R-HSA-212165          | Epigenetic regulation of gene expression                | −21.2| −18.6        | 18/149        |
| R-HSA-2559582         | Senescence-Associated Secretary Phenotype (SASP)        | −20.1| −17.6        | 16/112        |
| R-HSA-2559580         | Oxidative Stress Induced Senescence                     | −19.3| −16.8        | 16/125        |

**Downregulated:** Representative enriched BPs and pathways suppressed by SMG

| Term                  | Description                                             | LogP | Log(q-value) | InTerm_InList |
|-----------------------|---------------------------------------------------------|------|--------------|---------------|
| GO:0097352            | autophagosome maturation                                 | −6.1 | −2.6         | 4/46          |
| GO:0010038            | response to metal ion                                    | −5.9 | −2.6         | 7/352         |
| GO:0000422            | autophagy of mitochondrion                              | −5.2 | −2.1         | 4/75          |
| GO:0016049            | cell growth                                             | −5.0 | −2.0         | 7/473         |
| GO:0007005            | mitochondrion organization                              | −4.6 | −1.8         | 7/551         |
| GO:1901137            | carbohydrate derivative biosynthetic process            | −4.5 | −1.8         | 8/783         |
| GO:0040017            | positive regulation of locomotion                       | −4.3 | −1.7         | 7/607         |
| GO:0006695            | cholesterol biosynthetic process                        | −3.7 | −1.2         | 3/73          |
| GO:0042391            | regulation of membrane potential                         | −3.2 | −0.9         | 5/431         |

**KEGG Pathway**

| Term                  | Description                                             | LogP | Log(q-value) | InTerm_InList |
|-----------------------|---------------------------------------------------------|------|--------------|---------------|
| ko0520                | Amino sugar and nucleotide sugar metabolism             | −6.0 | −2.6         | 4/48          |

**Reactome Gene Sets**

| Term                  | Description                                             | LogP | Log(q-value) | InTerm_InList |
|-----------------------|---------------------------------------------------------|------|--------------|---------------|
| R-HSA-1474244         | Extracellular matrix organization                       | −3.9 | −1.4         | 5/301         |
| R-HSA-556833          | Metabolism of lipids                                    | −2.2 | −0.2         | 5/740         |
Next, mitochondrial membrane potential in SMG-CMs versus c-CMs was determined using the JC-1 staining. The JC-1 dye is a lipophilic, cationic dye that enters and accumulates into mitochondria to form so-called “J-aggregates”. Monomeric JC-1 emits green fluorescence, whereas J-aggregates present as red in functionally intact negatively charged mitochondria with a normal membrane potential. In contrast, in non-intact (damaged) mitochondria, JC-1 entry is greatly impeded due to loss of mitochondrial membrane potential resulting in a reduction in J-aggregates; then, monomeric JC-1 retains its green signal. As shown in Figure 3A, a decrease of the red fluorescence intensity was observed in SMG-CMs in parallel with increased green signal, suggesting the impairment (or full loss) of mitochondrial membrane potential (Figure 3A). As a consequence of mitochondrial potential dysfunctions, ATP synthesis is expected to be compromised. We
therefore determined ATP levels in SMG- and c-CMs using a luminescence-based assay. 48 h of SMG expo-
sure resulted in an almost 45% reduction of the ATP synthesis in SMG-CMs compared to controls (Figure 3C).

Finally, our Metascape analyses also identified changes in genes relevant to mitochondrial organization
(Table 1 and 2). We followed this up by investigating mitochondria morphology under SMG using TEM.
SMG exposure of CMs led to the hyperfusion of mitochondria, which were also morphologically different
compared to those in c-CMs, with SMG-CM mitochondria appearing ~40% longer than control ones (Figure 3D).
These findings further support our hypothesis that SMG exposure leads to impaired mitochondrial
turnover and therefore promotes a senescence-like phenotype in CMs.

SMG induces an increase of ROS, RNS, and stimulates the β-galactosidase activity in CMs
Our transcriptome analysis showed significant upregulation of genes involved in senescence and oxidative stress
pathways in SMG-CMs. To prove whether cellular ROS and RNS production is generally increased in SMG-CMs,
we next performed DHE (di-hydro-ethidium) stainings after 48 h of SMG exposure. In the absence of excessive
ROS, DHE remains cytosolic and emits blue fluorescence. Upon oxidation by free radicals, DHE is converted
into ethidium bromide that enters the nucleus and binds DNA to emit red fluorescence. SMG-CM nuclei present
with significantly increased levels of red signal (i.e., ~4-fold more red dots compared to c-CMs; Figure 4A). As
shown in Figure 4B, we could confirm the SMG-induced increase of ROS in IMR90- and iCell-CM by applying
the fluorescence DCFH-DA ROS assay. These findings confirm the accumulation of substantial ROS amounts
following 48 h exposure of CMs to SMG. The reactive nitrogen species (RNS) family of molecules is formed
from nitric oxide (NO) generated by the nitric oxide synthase (NOS) and superoxide anion (O_2^-) produced by
the NADPH oxidase, respectively. We also assessed RNS production in SMG-CMs, to identify additional reactive
species (Figure 4C). Last, to confirm senescence induction in SMG-CMs, cells were cultured in the presence of the

Figure 2. Exposure to SMG does not cause cytoskeletal deteriorations in CMs
(A) A representative immunofluorescence staining of sarcomeric α-actinin (ACTN2) and cardiac troponin T (cTnT) in c-CMs
and SMG-CMs (scale bar: 10 μm).
(B) TEM images of SMG-CMs and c-CMs. Black arrow heads indicate caveolae structures (black arrows show stress fibers
(scale bar: 500 nm).
(C) Western blot analysis of Caveolin (CAV1) in control and SMG-exposed CMs (molecular weight of 22 kDa). GAPDH
(36 kDa) was used as an internal protein loading control.

Figure 3. SMG reduces mitochondrial turnover and ATP levels
(A) A representative immunofluorescence staining of succinate dehydrogenase (SDH) in c-CMs
and SMG-CMs (scale bar: 10 μm).
(B) Graph showing the percentage of SDH-positive CMs in control and SMG-exposed CMs.
(C) Western blot analysis of oxidative phosphorylation enzymes in control and SMG-exposed CMs.

Figure 3. Effects of 48 h SMG on mitochondrial morphology, ROS production, membrane potential, and ATP synthesis

(A) Mitochondria staining of the SMG-CMs and c-CMs with the JC-1 dye to visualize disturbances in mitochondrial membrane potential (MMP). The JC-1 dye is a lipophilic, cationic dye that can be accumulated into the mitochondria to form the J-aggregates. The monomeric JC-1 naturally exhibits green fluorescence, whereas the JC-1 aggregates exhibit red fluorescence. Intact cells exhibit normal mitochondrial membrane potential. In this case, JC-1 dye enters and accumulates more in the negatively charged mitochondria and emits red fluorescent hence mitochondria appear redder as compared to mitochondria emitting green fluorescence. In contrast, JC-1 incorporation into mitochondria is greatly hindered due to the loss of mitochondrial membrane potential resulting in a reduction of J-aggregate formation retaining in its original green fluorescence. Both, monomers (green) and aggregates (red) were captured by using a confocal laser scanning microscope. The images were then processed and analyzed by using the ImageJ software. The ratio of green vs red color represents disturbances of the mitochondrial membrane potential; the greener, the intact mitochondria (scale bar = 10 μm).

(B) Fluorescence lifetime imaging microscopy (FLIM) of DyRed1-E5^-CMs (see video record 2 and DyRed1-E5^-SMG-CMs (see record 3) (Excitation length 488 nm). Visualization of newly synthesized and non-oxidized MitoTimer was performed setting an emission length of 497–531 nm (green color); Oxidized DyRed1-E5 was visualized at a fluorescence length of 583–695 nm. The ratio of green vs red fluorescence is representative for the overall mitochondrial turnover in the cells (scale bar: 20 μm).
SMG disrupts calcium flux in CMs

Our “omics” data pointed to several genes involved in calcium homeostasis and ion transport being deregulated in SMG-CMs, including G6PD, P2RX4, TSPO, ASPH, DMD, CACNA1C, NOL3, END1, CAV1, APP, CACR4, STC1, KCNA5, SLC38A1, RYR2, and ATP1A2. Therefore, we extended our studies to monitor Ca\(^{2+}\) flux in SMG—as compared to c-CMs. Cells were loaded with FLUO4-AM to monitor sarcoplasmic Ca\(^{2+}\) concentration ([Ca\(^{2+}\)]) during contractions. Transients were analyzed for the following kinetic parameters: time-to-peak (TTP, the time between the start and the peak (F\(_\text{max}\)) of the transient), (\(\Delta F/\Delta T\)\(_\text{max}\)) (the maximal slope within the TTP period), and T90% (the time for 90% decay from F\(_\text{max}\)). Exposure to SMG for 48 h had a significant impact on the kinetics of calcium transients in CMs, while no differences in sarcoplasmic resting [Ca\(^{2+}\)] were observed (Figures 5A). We did observe a significant reduction in (\(\Delta F/\Delta T\)\(_\text{max}\)) in SMG-CMs, indicating a slower rise in sarcoplasmic [Ca\(^{2+}\)] (Figure 5B). Accordingly, TTP displayed a significant increase (Figure 5C). Removal of Ca\(^{2+}\) from sarcoplasm was also slower in SMG-CMs, as indicated by the significantly longer T90% interval (Figure 5D). To monitor potential differences in the electrophysiology of Ca\(^{2+}\) currents between SMG-CMs and c-CMs, patch-clamp experiments were performed. Ca\(^{2+}\) currents were elicited by depolarization from the holding potential to potentials ranging from -60 to +60 mV and were normalized to the membrane capacitance (pA/pF, current density). As illustrated in Figure 5E, the pA/pF ratios in SMG-CMs were significantly lower than those recorded in c-CMs, indicative of significantly lower net sarcoplasmic Ca\(^{2+}\) entry following SMG exposure (Figure 5E). This finding indicates that SMG could impair normal L-type calcium channel (LTCC) function. Since RYR2 and phospholamban are critical elements for the kinetics of sarcoplasmic Ca\(^{2+}\) release and its reuptake into the sarcoplasmic reticulum, we tested whether the expression of the two proteins is affected in SMG-CMs. Western blot analysis revealed a reduction of the levels of both proteins (Figure 5F).

SMG affects the contraction and relaxation velocity of beating CMs

To investigate the effects of SMG on the contraction and relaxation velocity of CM sarcomeres, we generated one more transgenic iPSC line, ACTN2-eGFP+-hiPSCs (Figure S3), allowing the generation of ACTN2-eGFP+-CMs and real-time monitoring of several parameters characteristic of CM functionality. ACTN2 is a cardiac-specific cytoskeletal protein localizing to the Z-discs and playing a vital role in cardiac contractions. Using this line, we can identify ACTN2-rich Z-discs based on fluorescence in both control and SMG-CMs (Figures 6A and 6B). Figures 6C and 6D show representative measurements of the contraction and relaxation velocities recorded in c-CMs and SMG-CMs, respectively (data extracted from Videos S3 and S4, respectively, using the MATLAB-based SRAC-TRAC tool (Toepfer et al., 2019)). These recordings allowed detailed monitoring and analysis of contraction and relaxation of the individual z-discs. After analysis of the data generated from 6 independent experiments, we observed significant reduction in both contraction and relaxation velocities in SMG-CMs compared to c-CMs (Figures 6E and 6F). In addition, we also observed longer TTP duration in SMG-CMs while the contraction velocity slowed (Figure 6G), and the beating rate was significantly reduced (Figures 6H and 6I). In general, cardiomyocytes were responded to the adrenoreceptor agonist Isoprenaline (Video S5), and to the L-type agonist Bay-K8864 (Video S6) (see STAR Methods).

DISCUSSION

Human heart remodeling has been extensively studied under microgravity conditions and it is characterized by reduced oxygen metabolism and atrophy (Patel and Kos, 2005; Perhonen et al., 2001; Summers et al., 2005). Understanding the molecular mechanisms behind these phenotypes is of fundamental importance for developing preventive and therapeutic strategies against disease stemming from such environmental factors as microgravity and radiation. To this end, we used IMR90-hiPSCs-derived CMs to identify here a microgravity-controlled axis causing a senescence-like phenotype leading to...
contractile dysfunctions in CMs. To get evidence that SMG may induce also atrophy of the CMs, we investigated the effects of SMG on the expression of three genes MurF1, FOXO and SMAD3 that serve as markers for cardiomyocytes atrophy (Du et al., 2021; Ni et al., 2006; Tsui et al., 2015; Xin et al., 2017). As indicated in Figure S4 the, expression of all three genes was significantly upregulated in comparison to c-CMs.

Cellular senescence is one of the causal processes of aging. It invokes molecular cascades countering cell death when cells are exposed to various stress conditions. In the senescent state, cells are metabolically active, but no longer replicating (Payea et al., 2021). Senescence is also implicated in the development of aging-related disease (van Deursen, 2014). The main hallmarks for CM senescence are DNA damage response (DRS), mitochondria dysfunction, compromised contractile behavior, as well as hypertrophy (Tang et al., 2020). Our “omics” data analysis of genes and proteins differentially expressed following microgravity exposure suggested that SMG triggers mitochondrial-dependent senescence and modifies mitochondrial turnover (i.e., the adaptation of mitochondrial volume, content, and activity to current cellular energy needs) in CMs. This can be attributed to elevated oxidative stress levels, as ROS production can drive CMs to a senescent state, explain the imbalance between energy demand and supply (Ruiz-Meana et al., 2019), and lead to full-fledged heart disease (Kuster et al., 2010; Tian et al., 2021).

Figure 4. Effects of SMG on ROS, RNS production, and β-galactosidase activity in CMs
(A) Fluorescent images showing an increase in DHE staining in SMG-CMs in comparison to c-CMs. Red staining in nuclei indicates a net increase in the ROS production (Nuclei: DAPI) (Scale bar: 20 μm). Quantification was performed using ImageJ software (red: nuclear DHE, blue: DAPI). Five images of c-CMs (in total n = 71 cells) and SMG-CMs (in total n = 53 cells) from tree independent experiments (3 slides) were selected and all the cells in each image were counted for positive and negative red nuclear staining. Then, the percent positive staining was calculated and represented in the graph as % DHE staining (Mean ± SD, p < 0.05).

(B) ROS quantification in c-CMs and SMG-CMs by the fluorescence DCFH-DA test. The fluorescence intensity was measured using Softmax Pro M5e 96-well plate reader with excitation and emission wavelengths of 485 and 535 nm, respectively. All the fluorescence intensities are normalized with control c-CMs (100%) (Mean ± SD, p < 0.05, n = 9, three independent experiments, performed in triplicates).

(C) Reactive nitrogen species (RNS) production in SMG-CMs and c-CMs. Fluorescence was quantified in a fluorometric plate reader at 480 nm/530 nm (expressed as percent of the absolute values of RNS in c-CMs (mean ± SD, p < 0.05, n = 9, two-tailed t-test; three independent experiments, performed in triplicates).

(D) Bright filter and fluorescent images showing increase β-galactosidase staining in SMG-CMs and c-CMs (Scale bar = 50 μm) in IMR90-and iCell-CMs. Quantification of the GFP intensity from 35 CMs from each condition (c-CMs and SMG-CMs, 3 independent experiments) was determined by the ImageJ software. Then, the raw intensity data were plotted as a box plot.
ROS is available in at least eight cellular compartments, namely in mitochondria, lysosomes, endoplasmic reticulum, the Golgi, peroxisomes, the nucleus, and in the cytosol (Naviaux, 2012; Zorov et al., 2014). Cytosolic ROS could trigger complex cellular signaling pathways that induce ROS release from mitochondria, a process known as the ROS-induced ROS release mechanism (RIRR) (for review see (Zorov et al., 2014)). Indeed, we demonstrated that SMG caused a significant increase in cellular ROS. However, further studies are required to demonstrate whether the RIRR mechanism is involved in the SMG-induced increase of the ROS level. As indicated in Figure S4, the expression of the antioxidant genes, superoxide dismutase 2, mitochondrial (SOD2), and catalase (CAT) was upregulated in the SMG-CMs. The upregulation of the antioxidant enzyme SOD2 may occur due to a potential inactivation of the SMG-induced increase of intracellular ROS thereby preventing the senescent CMs from cell death. However, this hypothesis should be underpinned by extensive enzymatic and other assays.

We also examined the effect of the SMG on the cell viability of the IMR90 CMs and iCell CMs. As shown in Figure S5, the cell viability of the CMs which were exposed to 48 h SMG was approximately 85%, whereas further decreased by 74% after 72 h SMG. Therefore, we suggest that a 48 h SMG duration time is an optimal period for our SMG-related investigations. Our viability data suggest that during 48 h SMG most of the cells are in the senescent stage avoiding an extensive cell death that can happen extensively due to a syndrome of stress signals, e.g., due to starvation of the cells. Moreover, during the 48 h SMG period, no exchange of the medium is required. A SMG period of more than 48 h would necessitate medium exchange under 1 g conditions and therefore would lead to inconsistent findings. Interestingly, we identified by our Metascape analysis that 27 genes belonging to the p53 pathway and 8 genes belonging to TOR pathway whose expression was suppressed in SMG-CMs (see Table S3 and Table S7A). Normally, these pathways are involved in cell death processes but senescent cells, in general, are resistant to apoptosis and cell death (for review see (Hu et al., 2022)). Therefore, we may speculate
that the SMG-CMs suppressed genes associated with these pathways preventing senescent CMs from apoptosis and cell death.

In our study system, this is evident in the effects SMG has on the contractile function of CMs. By challenging intracellular Ca\(^{2+}\) homeostasis, SMG led to rhythmic changes in the contraction and relaxation times of CMs. The increase in contraction velocity slopes recorded in SMG-CMs is indicative of SMG acting as

**Figure 6. Live imaging of contraction and relaxation velocity activity of ACTN2-eGFP\(^{-}\)-CMs and ACTN2-eGFP\(^{-}\)-SMG-CMs**

(A and B) Fluorescence microscopy shows the Z-discs of ACTN2-eGFP\(^{-}\)-CMs in c-CMs and ACTN2-eGFP\(^{-}\)-SMG-CMs (ACTN2 is enriched in Z-discs) immediately after 48 h SMG. ACTN2 Video record 4 and 5 shows the contractile activity of the c-CMs and SMG-CMs, respectively. Fluorescence microscopy has been performed using the Olympus FluoView1000 confocal system.

C–D, Contractile velocity of c-CMs and SMG-CMs (representative experiments; see Videos S3 and S4, respectively) was determined by analyzing the video records 4 and 5 (50fps), respectively by using the VA1.9 software. E–G, Diagrams show the duration of contraction, the duration of relaxation, and the TTP values and of the c-CMs and SMG-CMs, respectively (mean ± SD, n = 6, *p < 0.05; two-tailed t-test; 3 independent experiments). H, Diagram shows the \(\frac{\Delta V}{\Delta T})_{\text{max}}\) ratio in SMG-CMs versus c-CMs (mean ± SD, n = 6, *p < 0.05; two-tailed t-test; 6 independent experiments). I, beating frequency of SMG-CMs versus c-CMs (mean ± SD, n = 11, *p < 0.05; 3, two-tailed t-test; independent experiments).
negative inotropic factor. This likely manifests as a result of cell membrane destabilization, as evidenced by the structural deformities and caveolae formation observed in SMG-CMs. In addition, the higher $T_{90}$ values in SMG-CMs suggest a delayed $Ca^{2+}$ removal from the sarcoplasm of these cells. Such a change in intracellular $Ca^{2+}$ clearance rates has also been observed in aged mice cardiomyocytes (Yang et al., 2006) and may be justified by an impaired function of SERCA2a. We have demonstrated that the $Ca^{2+}$ influx was significantly lower in SMG-CMs (Figure 5E). On the other hand, the expression of several calcium ion homeostasis genes (Table S2 and Table S7B) was upregulated although cardiac muscle contraction-specific genes were significantly downregulated (Table S3 and Table S7C). From these findings, we may speculate that the L-type calcium channel (LTCC) activity is impaired in SMG-CMs via biomechanical mechanisms.

Finally, the onset of senescence has been shown to confer dramatic changes to 3D genome organization, including reshuffling of TADs (Zirkel et al., 2018) and chromatin compaction (Criscone et al., 2016). Given that gene expression changes induced by SMG resembled those seen in replicative senescence, we used iHi-C and could show that similar changes occur in CMs under SMG conditions. On the one hand, we recorded strengthened longer-range interactions at the expense of shorter-range ones, which can be indicative of chromatin compaction. But, in contrast to what occurs during replicative senescence, we find that the majority of TADs that change their boundaries due to SMG appear to split into smaller domains ([in senescence the major effect is TAD merging; (Zirkel et al., 2018)]. This difference could well be due to the post-mitotic nature of CMs or a signature of SMG-induced genome reorganization.

More recently, human Jurkat T cells were exposed to short-term gravitational changes in parabolic flight and suborbital ballistic rocket flight experiments (Vahlensieck et al., 2021). Authors reported that smaller chromosomes (Chr 16–22 except for chr18) have higher intrachromosomal interactions compared to large chromosomes in altered gravity. However, it should be pointed out, that during the parabolic flight, the Jurkat T cells were exposed to 31 alternating hypergravity, microgravity, and hypergravity phases, each lasting 20–22 sec (Acharya et al., 2019). In striking contrast, in the present study, CMs were continuously exposed exclusively to microgravity for 48 h. Therefore, due to fundamental experimental differences of the Hi-C data in our study and the study reported (Vahlensieck et al., 2021) are not surprising. Nevertheless, it is evident that SMG also impacts large-scale chromosomal organization.

However, to validate our key findings, we also repeated some key experiments with the commercially available iCell-CMs. The iCell-CMs were derived from hiPSCs after reprogramming blood cells (according to the declarations of the Cellular Dynamics). Again, SMG induced an elevation of intracellular ROS level, an increase of the $\beta$-galactosidase activity and significantly reduced the beating frequency of the iCell-CMs. Also, similar effects of SMG were observed regarding the viability of the IMR90-CMs and iCell-CMs. However, the comparative experiments were performed with purified cardiomyocytes in the absence of other non-myocyte cells such as cardiac fibroblasts which may affect the function of cardiomyocytes. This limitation can be improved by investigating the effects of SMG on 3D cardiac organoids consisting of cardiomyocytes and non-myocyte cells such as fibroblasts and endothelial cells.

Overall, our observations are consistent with the impairment of CM contractile function owing to challenges in mitochondrial metabolism, which is a fundamental aspect of cardiac aging (Barton et al., 2017). Taken together, despite some limitations, our results can lay the foundation for developing strategies to prevent senescence processes in the heart and to develop new therapeutics against microgravity-induced disease.

**Limitations of the study**

There are three limitations of the present study. One important limitation is the use of the hiPSC-derived cardiomyocytes which till now are not mature although they are fully functional. The other limitation is the short time of 48 h exposure under simulated microgravity conditions. Indeed, experiments under real microgravity conditions onboard the International Space Station (ISS) for a longer period would be optimal. Finally, further experiments are needed to demonstrate whether the senescence processes are reversible in the presence and absence of anti-senescence compounds.

**STAR★METHODS**

Detailed methods are provided in the online version of this paper and include the following:

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QUANTIFICATION AND STATISTICAL ANALYSIS

SUPPLEMENTAL INFORMATION
Supplemental information can be found online at https://doi.org/10.1016/j.isci.2022.104577.

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AUTHOR CONTRIBUTIONS
A.A. conducted the main experiments, collected all the findings, and contributed to the writing of the manuscript. K.P. and K.K. contributed to the experimental part. H.N. generated the plasmids for the study. S.P. conducted the Ca²⁺ measurements. F.N., J.H., and T.S. conducted and supervised the patch-clamp experiments. R.H. supervised the simulated microgravity experiments. A.M. conducted iHi-C experiments and E.G.G. the bioinformatic analysis of the iHi-C data. A.P. contributed to the analysis and interpretation of data, and to the writing of the manuscript. A.S. designed and supervised the study, interpreted data, conducted analysis of “omics” data, and wrote the manuscript.

DECLARATION OF INTERESTS
The authors declare no competing interests.

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## STAR METHODS

### KEY RESOURCES TABLE

| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| **Antibodies**      |        |            |
| Anti-Caveolins-1    | ab18199 | Abcam      |
| anti RYR2           | ab219798| Abcam      |
| anti GAPDH          | AB516   | Sigma      |
| anti PLN            | MA3-922 | Thermo Fisher Scientific |
| anti-cardiac Troponin T | MA5-12960 | Thermo Fisher Scientific |
| **Bacterial and virus strains** | | |
| Stellar Competent Cells | 636766 | TAKARA Bio |
| One Shot™ TOP10 Chemically Competent E. coli | C404010 | Thermo Fisher Scientific |
| **Chemicals, peptides, and recombinant proteins** | | |
| Dihydroethidium (DHE), Blue fluorescent dye | ab145360 | Abcam |
| L-Arginine-HCl      | 89989   | Thermo Fisher Scientific |
| L-Lysine-2HCl       | 89987   | Thermo Fisher Scientific |
| L-Arginine-HCl, 13C6, 15N4 | 89990 | Thermo Fisher Scientific |
| L-Lysine-2HCl, 13C6, 15N2 | 88209 | Thermo Fisher Scientific |
| **Critical commercial assays** | | |
| Applied Biosystems™ SYBR™ Green PCR Master Mix | 43-444-63 | Applied Biosystems™ |
| ATPlite Luminescence Assay System | 6016943 | Perkin Elmer |
| β-galactosidase staining kit | C10850 | Thermo Fisher Scientific |
| Oxiselect Intracellular ROS Assay Kit (Green Fluorescence) | MBS168048 | My Biosource |
| MitoProbe™ JC-1 Assay Kit | M34152/22204 | Thermo Fisher Scientific |
| **Deposited data** | | |
| RNAseq transcriptome data | This paper | ArrayExpress, EMBL-EBI Datasets platform: E-MTAB-11159 |
| iHC2.0 data | Mizi et al. (2020), Methods 170, 33-37 | https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE161636 |
| **Experimental models: Cell lines** | | |
| IMR90 hiPSCs | iPS(IMR90)-4-DL-1 p18+34(9) | Wisconsin International Stem Cell Bank |
| iCell Cardiomyocytes | iCell Cardiomyocytes, 11713 | Cellular Dynamics, Fujifilm |
| **Oligonucleotides** | | |
| Primers for qRT-PCR for several genes | See Table S8 | Integrated DNA Technology (IDT), Integrated DNA Technology, Leuven, Belgium |
| **Recombinant DNA** | | |
| pMitoTimer | 52659 | Addgene |
| pCAG-PBase | 40972 | Addgene |
| PB-TA-ERP2 | 80477 | Addgene |
| pCP-CMV-GCaMP6s | 40753 | Addgene |
| PB-TAC-ERP2 | 80478 | Addgene |
| cACTN2 tagging | N-Terminal CopGFP tagging | CRISPR/Alstem Bio |
| IMR90-ACTN2-copGFP | Custom made |

(Continued on next page)
RESOURCE AVAILABILITY

Lead contact

Further information and requests for resources and reagents should be directed to and will be fulfilled by the lead contact, Agapios Sachinidis (a.sachinidis@uni-koeln.de).

Materials availability

All reagents will be made available. The following cell lines will be available after completion of a Materials Transfer Agreement.

- A transgenic ACTN2-eGFP+-hiPSCs line (IMR90).
- A transgenic Tet-inducible MitoTimer hiPSCs line (IMR90).
- IMR90-ACTN2-copGFP (recombinant DNA).

Data and code availability

- The RNAseq transcriptome data were deposited on the ArrayExpress EMBL-EBI Datasets platform (accession number: E-MTAB-11159).
- The iHi-C 2.0 data were deposited on NCBI, GEO (accession number GSE16163).
- The paper does not report original codes and reagents. All codes and reagents used in this study are listed in the key resources table. Any additional information required to reanalyze the data reported in this paper is available from the lead contact upon request.

METHOD DETAILS

Differentiation of hiPSC into cardiomyocytes

Experiments were performed with the IMR90 hiPSC (authorized by the Robert-Koch Institute; Berlin, Germany, license number: AZ 3.04.0210083). Cells were cultured on matrigel-coated Petri dishes. Cells were cultured in StemMACS™ iPS-Brew XF media (Milteny Biotech, Germany) supplemented with 50 U/mL penicillin and 50 U/mL streptomycin (Thermo Fisher Scientific, MA, USA), at 37°C and 5% CO2. After reaching of a confluence of 90%, cells were dissociated by trypsinization using the trypLE solution (Thermo Fisher Scientific, MA, USA) and propagation of the cells was performed in StemMACS™ iPS-Brew XF media in the presence of 10 μM ROCK inhibitor. Differentiation of hiPSCs to cardiomyocytes was performed by a combination of two differentiation protocols (Lian et al., 2012; Tohyama et al., 2013) as we described previously (Nemade et al., 2020). The cells were cultured in RPMI1640/B-27−insulin medium (Medium without insulin (B-27−insulin medium)) in the presence of 10 μM CHIR99021 (Tocris, United Kingdom) for 24 h (day 1). Then the culture medium was replaced with a normal RPMI/B-27−insulin medium and cells were cultured for further 24 h (day 2). Then the culture medium was replaced with RPMI/B-27−insulin media containing 10 μM IWP2 (Tocris, United Kingdom) and cells were cultured for a further 48 h (day 4). Then the medium was replaced with basal RPMI/B-27−insulin medium and was refreshed every 2 days. Spontaneous beating cardiomyocytes were observed from day 8 onwards. Enrichment of the cardiomyocytes was performed by culturing the cells with glycode-free RPMI (Thermo Fisher Scientific, MA, US) containing 4 μM sodium DL-lactate for 6 days. The purity of cardiomyocytes at day 18 of

| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| ImageJ for fluorescent image analysis | (Crowe and Yue, 2019; O’Brien et al., 2016) | https://imagej.nih.gov/ij/ |
| Video Analyzer 1.9 | GitHub software repository; Acharya et al. (2022); 11.1280. https://doi.org/10.3390/cells11081280 | https://github.com/nblackburn123/Video-Analyser/releases/tag/v1.9 |
| Maxquant (version 1.5.3.8) | Nat Protoc 11, 2301–2319. | https://www.maxquant.org/ |
| iHiC2.0 | Mizi et al. (2020); Methods 170, 33-37 | https://www.sciencedirect.com/science/article/abs/pii/S1046202319300106 |
For all the studies, the cells from the central 3 mm area, which were exposed to microgravity of 0.012 g range from 0.012 to 0.001 g. To remove all the air bubbles, Rotation of flask generates acceleration, which exposes cells to SMG in the 2D clinostat. In the Slide-flask 2D-clinostat, the fast and constant rotation at 60 rpm induces gravity vector control CMs were cultured in slide-customized flasks and were kept in the incubator but not placed on the 2D clinostat. In the Slide-flask 2D-clinostat, the fast and constant rotation at 60 rpm induces gravity vector randomization and during this rotation, to avoid any fluid movements inside the flasks, it is necessary to remove all the air bubbles. Rotation of flask generates acceleration, which exposes cells to SMG in the range from 0.012 g (at the 3 mm area from the center of rotation) to 0.036 g at the farther distance. For all the studies, the cells from the central 3 mm area, which were exposed to microgravity of 0.012 g for different times and were harvested for further experiments.

MTT assay

The IMR90-CMs and iCell-CMs were seeded in a single channel ibidi slide and incubated in the cell culture incubator for 4 days. The cells were then exposed to SMG on 2D-Clinostat. Post 24 h, 48 and 72 h of exposure, 200 μL MTT solution was added and cells were incubated in the dark for 150 min. Then the medium was removed and formazan crystals formed in the cells were dissolved using 200 μL of DMSO followed by transfer into a 96 well plate. The absorbance was read at 570 nm using Softmax Pro M5e 96-well plate reader (Molecular Devices, Sunnyvale, CA, USA) (n = 9; three independent experiments) and represented as per cent cell death compared to that of control cells.

RNA isolation from CMs

RNA isolation was done by using RNAqueous™ Micro Total RNA Isolation Kit AM1931 from (Ambion). Total RNA concentration was measured by UV-Vis spectrophotometer nanodrop2000c (Thermo Fisher Scientific). Furthermore, samples were measured on the Fragment Analyzer 5200 (Agilent). In case of gDNA isolation from CMs, RNA isolation was done by using RNAqueous Micro Total RNA Isolation Kit AM1931 from (Ambion). Total RNA concentration was measured by UV-Vis spectrophotometer nanodrop2000c (Thermo Fisher Scientific). Furthermore, samples were measured on the Fragment Analyzer 5200 (Agilent). In case of gDNA
contamination, the DNase I (Thermo Fisher Scientific) digest was performed. Then samples were processed for library preparation for RNA seq.

**RNAseq library preparation**

Sequencing-ready libraries were produced using a QuantSeq 3’ mRNA-Seq Library Prep Kit FWD for Illumina (015UG009V0241) following the standard procedures, as described by Lexogen. Indexed library preparation was performed to allow multiplexed sequencing. 250 ng mRNA samples were used as input for the library preparation. Finished libraries were quality controlled on a Bioanalyzer device (Agilent), using the HS-DNA assay. The concentration of obtained libraries was quantified using a Qubit dsDNA HS assay (Thermo Fisher). A sequencing-ready pool of indexed libraries was prepared according to these quantifications.

**Sequencing of prepared library**

Samples were pooled in equimolar ratio and the library pool was quantified using a Qubit dsDNA HS assay kit (Thermo Fisher Scientific). Sequencing was performed on Illumina NextSeq 500 with a SR75 High Output by following Lexogen standard protocol. The transcriptome RNA-seq data were validated by qRT-PCR analysis of upregulated (Figure S7A) and downregulated genes (Figure S7B) (for prime sequence see Table S8).

**SILAC labelling and mass spectroscopy**

We performed quantitative proteomics applying the “stable isotope labeling by/with amino acids in cell culture” (SILAC) methodology as described previously (Chen et al., 2015) to identify proteins dysregulated in under microgravity conditions in CMs. For this aim, hiPSCs were cultured with normal RPMI 1640 containing “heavy” and “light” arginine and lysine isotopes of both amino acids and supplemented with GlutaMAX (1%, v/v). The “heavy” SILAC RPMI medium (500 mL) was supplemented with 50 mg of 1-Arg-10 = L-[13C615N4] Arg. HCl, 25 mg of 2-Lys-8 = L-[13C615N2] Lys.HCl, 100 mg proline and 5 mL pen-strep. The composition of the “light” SILAC RPMI 1640 medium (500 mL) was similar but the heavy amino acids were replaced with 50 mg light 1-Arg-0 = l-[12C614N4]Arg.HCl and 25 mg light 2-Lys-0 = l-[12C614N2] Lys.HCl and 100 mg proline and 5 mL pen-strep. The hiPSCs were cultured in parallel in “heavy” and “light” SILAC media till the cells reached a confluence of more than 90% (day 0). Then cells were differentiated in SILAC media in the absence and presence of CHIR99021 (10 μM) and IWP-2 (5 μM) and 4 mM L-lactate as described by the differentiation protocol. In total after 20 days of differentiation, the “heavy” and “light” labelled CMs were further cultured in a maturation medium containing the “heavy” or the “light” amino acids at the same concentrations as shown above. The SILAC labelled experiments were divided into two groups; the “heavy” and the “light” isotope group. The “light” isotope labelled cells were exposed to SMG and the “heavy” isotope labelled cells were kept as ground control. After exposing the cells to SMG, CMs cells from both groups were lysed using standard RIPA lysis buffer (Sigma). Then “heavy” and “light” samples were mixed at a ratio of 1:1 and subjected to single-pot, solid-phase-enhanced sample-preparation (SP3) digestion following mass spectrometry analysis. All samples were analysed on a Q-Exactive Plus (Thermo Scientific) mass spectrometer that was coupled to an EASY nLC 1000 UPLC (Thermo Scientific) chromatograph. Peptides were loaded with solvent A (0.1% formic acid in water) onto an in-house packed analytical column (50 cm × 75 μm I.D., filled with 2.7 μm Poroshell EC120 C18, Agilent) and chromatographically separated at a constant flow rate of 250 nL/min using 60 or 150 min gradient of 5-30% solvent B (0.1% formic acid in 80% acetonitrile). The mass spectrometer was operated in data-dependent acquisition mode. The MS1 survey scan was acquired from 300-1750 m/z at a resolution of 70,000. The top 10 most abundant peptides were isolated within a 2 Da window and subjected to HCD fragmentation at a normalized collision energy of 27%. Precursors were dynamically excluded for 20 s. For bioinformatics data analysis all raw data from mass spectrometry were processed with Maxquant (version 1.5.3.8) using default parameters (Tyanova et al., 2016). Protein SILAC ratios were log2 transformed and normalized to the median. For every time point, a one-sample t-test was performed. Proteins with a p value < 0.05 and log2 change < -1 or >1 were considered significantly changing and subjected to profile clustering, which was generated using the z-scored mean values per group. A two-sample t-test was used to determine significantly enriched proteins. Potential contaminants were annotated based on the Crapome repository (Mellacheruvu et al., 2013).

**Quantitative real-time PCR for gene expression analysis**

The mRNA expression analysis was executed as we described previously (Nemade et al., 2020). In brief, 500 ng of mRNA was used to synthesize cDNA (high-capacity cDNA Synthesis Kit, qScript and Applied
bio-system). Then cDNA synthesis was diluted with highly pure RNAse free water at 1:5 ratio and then 2 μL of the diluted cDNA was used for qRT-PCR (Applied Biosystems 7500 FAST Real-Time PCR System) with the appropriate primers. GAPDH mRNA was used as an internal control.

Native whole-genome chromosome conformation capture (iHi-C 2.0) and data analysis

The effect of microgravity on the chromosome architecture in the CMs was investigated by the iHi-C 2.0 method as we described previously (Mizi et al., 2020). Briefly, CMs (0.3 × 10^6) were seeded on fibronectin-coated 12 cm² slide-flask and exposed to SMG for 48 h. Control CMs grown at 1 g were used as control. After SMG exposure cells were gently removed from the slide and processed for iHi-C2.0. First, the nuclei were isolated for digestion and ligation in naïve condition, followed by purification and linearization for DNA library preparations. We initiated the library preparation by adding 10 bp unique molecular identifier (UMI) oligo provided with the Select High Sensitivity kit (Agilent). Then we performed the paired-end sequencing, adapter primer was used into the barcode library, while utilizing the TruSeq SR cluster kit (Illumina). It reads up to 15 × 10^7 read pairs each and each read length was 75 nt. Raw sequencing reads in each pair (.fastq files) were mapped independently to the human reference genome (hg19) using BWA-MEM after trimming of 5 nt from each end to eliminate identical NlaIII sequence motifs. PCR duplicates were defined as those read pairs mapping to the same locations in the genome and carrying the same UMI, and were manually removed. Following this, reads were further filtered to remove any that did not precisely map next to genomic NlaIII recognition sites. Similarly, read pairs with both ends mapping to the same NlaIII fragment or with “dangling” ends were also removed. The resulting data matrices were processed using “Bloom” for imputation to counter sparsity (Gusmao et al., 2020). Ultimately, the library of filtered mapped read pairs were normalized and converted to.hic files and visualized via Juicebox (Durand et al., 2016) to give rise to KR-normalised (Knight and Ruiz, 2012), interactive, and zoomable iHi-C maps. Compartment analysis was performed using the first principal component of the Hi-C matrices as described (Rao et al., 2017) and TAD boundaries were identified using default setting in (Kruske et al., 2016). Binarization and matrix subtractions were performed as previously described (Zirkel et al., 2018).

ATP assay

Intracellular ATP levels in CMs were determined using the ATPlite Luminescence Assay System (PerkinElmer, Waltham, MA) according to the manufacturer’s instructions and the light intensity at 560 nm was quantified using the Softmax Pro M5e 96-well plate reader (Molecular Devices, Sunnyvale, CA, USA). The raw intensity data were normalized and presented as per cent changes of the ATP levels. Three independent experiments were performed in triplicates (n = 9).

Immunostaining (ICC)

CMs sarcomere assembly was detected by immunostaining of the cardiac-specific proteins ACTN2 and cardiac troponin T (cTnT). Cardiomyocytes were fixed by using ice-cold 99% methanol. Then cells were permeabilized with 0.3% TritonX-100 in the presence of 5% BSA. Then cells were washed and incubated with 1:200 diluted primary antibodies against α-actinin and cTnT at 37°C for 1 h. After 3 times washing with PBST, staining of the proteins was performed by incubating the cells for 1 h at RT with a secondary species-specific Alexa Fluor-488/568-conjugated secondary antibody (1:2000 dilution; Invitrogen, Darmstadt, Germany). The cells were washed and mounted on a slide with Prolong Gold anti-fade mount with DAPI (Invitrogen, Darmstadt, Germany). Fluorescence microscopy was performed with the Axiovert 200 fluorescence microscope and images were then analyzed using Axiosvision 4.3 software (Carl Zeiss).

Measurement of intracellular ROS in CMs post SMG exposure

The intracellular ROS production in CMs was determined by dihydroethidium (DHE) staining as described previously (Nemade et al., 2018). Briefly, cell staining was performed by culturing the cells with 5 μM DHE for 30 min at 37°C and 5% CO₂. Then cells were washed with DPBS (Ca²⁺ and Mg²⁺ free PBS; Thermo Fisher Scientific) and fixed with Prolong® Gold anti-fade mount with DAPI (Invitrogen, Darmstadt, Germany). Fluorescence microscopy was performed with the Axiovert 200 fluorescence microscope and fluorescence was quantified using Axiosvision 4.3 software (Carl Zeiss). The method is based on the oxidation of DHE by ROS to form ethidium bromide, which penetrates the nucleus and bind to DNA and due to red
punctate patterns appearing inside the nucleus. Analysis of arbitrary images has been executed by image J as described previously (Huang et al., 2019). Five images of c-CMs (in total n = 71 cells) and SMG-CMs (in total n = 53 cells) from three independent experiments (3 slides) were selected and all the cells in each image were counted for positive and negative red nuclear staining. Then the percent positive staining was calculated and represented in the graph as % DHE staining. Additionally, we determined the intracellular level of ROS, by using the 20, 70 dichlorofluorescin diacetate (DCFH-DA) test according to the manufacturer’s instructions (ABIN2344999, Cell Biolabs, San Diego, CA, USA). After 48 h of SMG, the cells were washed with PBS and incubated with DCFH-DA (10 μM) for 30 min at 37°C. DCFH-DA diffuses into cells and is deacetylated by cellular esterase to non-fluorescent DCFH, which is rapidly oxidized to highly fluorescent DCF in presence of ROS. The fluorescence intensity was measured using Softmax Pro M5e 96-well plate reader (Molecular Devices, Sunnyvale, CA, USA) with excitation and emission wavelengths of 485 and 535 nm, respectively (n = 9, three independent experiments, performed in triplicates). All the fluorescence intensities are normalized with control cells (100%) and represented as a percentage of the control.

Determination of the NO-derived-reactive nitrogen species (RNS)
The NO-derived RNS were quantified with the OxiSelect Intracellular Nitric Oxide Assay Kit following the manufacturer’s instructions (Cell Biolabs, Inc., San Diego, CA) (Novo and Parola, 2008). Experiments were performed in three independent experiments, performed in triplicates (n = 9). Fluorescence was quantified in a fluorometric plate reader at 480 nm/530 nm (expressed as percent of the absolute values of RNS in c-CMs).

Transmission Electron Microscopy of CMs
The TEM experiments were performed as we described previously (Nemade et al., 2018). Briefly, 48 h SMG exposed CMs and control CMs, were seeded into specially designed slide flask having 3 tier detachable ACLAR® embedding films (Ted Pella, Inc. USA) which were glued to slide flask shown in (Figure S1). The ultrathin slides (~70 nm) were prepared by using Leica EM UC6 Ultramicrotome (Leica, Germany). Then slices were subjected to dual stain with uranyl acetate followed by lead citrate. TEM imaging analysis was performed using a transmission electron microscope (EM109, Zeiss, Germany) equipped with a frame-transfer-CCD camera. Using ImageJ, the mitochondrial length was determined and plotted in a box plot.

Calcium imaging of CMs post SMG exposure
CMs (0.3 × 10^6) were seeded on fibronectin-coated 12 cm² slide-flasks and were exposed to SMG for 48 h. Cells were then loaded with 2 μM FLUO-4 AM (Invitrogen) for 30 min at 37°C and 5% CO₂, and thereafter were washed with basal media. Local [Ca²⁺] oscillations were then recorded by applying the line scan mode of the Olympus FV1000 microscope using a 60× oil immersion objective and an excitation wavelength of 488 nm, as described previously (Walter et al., 2016). Upon capturing, background-correction was applied to all transients For both SMG and c-CMs, we recorded seventy independent readings to perform a robust statistical analysis (n = 70, from 2 independent cell preparations).

β-galactosidase staining
To detect the activation of β-Galactosidase as a parameter for senescence processes in CMs the β-Galactosidase activity Cell Event™ Senescence Green Detection Kit (Thermo Fisher Scientific). The staining was performed at 37°C in the absence of CO₂ for 1 to 2 h. The enzyme cleavage the substrate β-galactosidase into monosaccharides in an acidic pH medium. A fluorescence product covalently interacts with intracellular proteins. Quantification of the fluorescence intensity was performed at absorption 490 nm and emission wavelength of 514 nm (three independent experiments, n = 35 CMs from each condition) was determined by the imageJ software. Then the captured images were evaluated by imageJ to analyse the senescence versus non-senescent CMs. CMs from each condition (c-CMs and SMG-CMs) were selected and the GFP intensity was quantified using the imageJ software. Then the raw intensity data were plotted.

Mitochondrial membrane potential (mΔψ) assay
Mitochondrial membrane potential differences in CMs were determined by using JC-1 dye staining. The CMs (0.3 × 10^6) were seeded onto fibronectin-coated 12 cm² slide-flask. After 72 h incubation, the cells
were exposed to SMG for the next 48 h by keeping respective ground control cells. Then cells were loaded with 10 μM JC-1 dye (AAT Bioquest, CA) and incubated for 30 min at 37°C, 5% CO₂. After incubation, the cells were washed two times with DPBS. H₂O₂ (100 nM; 48 h) was used as a positive control. After washing, cells were cultured in normal media and then the cells were subjected to microscopy.

**Protein expression analysis of SMG exposed CMs by Western blot**

The SMG exposed and in parallel, the ground control cells from the central 3 mm area of the flask were scratched and transferred into a 15 mL tube with 5 mL ice-cold PBS. After centrifugation at 1500 rpm for 5 min protein was extracted by bath sonication using RIPA lysis buffer (50 mM Tris-HCl pH 7.6, 150 mM NaCl, 1% TritonX-100, 1% sodium deoxycholate, 0.1% SDS; Sigma). Isolated protein was then quantified by using a standard BCA reagent kit (Thermo Fisher Scientific). Then protein samples were separated by 20% SDS-polyacrylamide gel electrophoresis. After electrophoresis proteins were transferred into the nitrocellulose membrane by using blot, a dry blot instrument (Thermo Fisher Scientific) and the transfer buffer. After the transfer, membranes were treated with 5% non-fat milk in TBS buffer for 1 h at room temperature. Membranes were then treated overnight with the primary antibodies in a dilution of 1:2000 to label the targeted protein. Membranes were then washed 3 times with TBST. The dyes IRDye 680CW and 800CW (Thermo Fisher Scientific) were used. Membranes were then incubated with the secondary antibody (1:12000 dilution) for 1 h. Then membranes were washed again for 3 times and then protein bands were detected by using the Odyssey Infrared Imaging System (LI-COR).

**Whole-cell patch-clamp of CMs post SMG exposure**

Electrophysiological recordings were performed using the whole-cell configuration of the patch-clamp technique as described previously (Hamill et al., 1981). Patch pipettes were prepared from thick-walled borosilicate glass capillaries (1.5/0.84 mm OD/ID; World Precision Instruments) using a P97 Micropipette puller (Sutter Instruments) and had resistances between 5 and 7 MΩ when filled with the Cs-based internal solution. The bath was connected to ground via 120 mM sodium chloride agar bridges. Currents were acquired at 50 kHz and filtered at 10 kHz using an EPC9 amplifier (HEKA, Germany) controlled with HEKAs Pulse software. Leak and capacitive currents were subtracted online by use of a –P/5 protocol and series resistance was compensated electronically by up to 80%. All experiments were performed at room temperature and from a holding potential of –80 mV. During the recordings, cells (n = 8 for c-CMs and n = 6 for SMG-CMs) were continuously superfused with external solution containing (in mM) 120 NaCl, 5 KCl, 3.6 CaCl₂, 1 MgCl₂, 10 HEPES, 20 tetrathylammonium chloride and 2 4-aminopyridine, with the pH adjusted to 7.4 using NaOH. The pipette solution contained (in mM) 120 CsCl, 3 MgCl₂, 10 ethylene glycol-bis(2-aminoethylether)-N,N,N',N'-tetraacetic acid (EGTA), 5 HEPES and 4 MgATP, with the pH adjusted to 7.3 using CsOH. Ca²⁺ currents were recorded in response to 100 ms voltage-steps after a pre-pulse at –40 mV to inactivate Na⁺ channels and normalized by the whole-cell capacitance to give current density values. To construct current-voltage-relationships, peak current densities measured between –50 and 60 mV (10 mV increments) were plotted as a function of the test potential and fitted with a combined Ohm-Boltzmann equation.

**Real-time imaging of mitochondrial ROS by MitoTimer**

To monitor the mitochondrial ROS a transgenic hiPSCs line (IMR90) was generated allowing the expression of the genetically encoded sensor protein MitoTimer in a Tet-inducible manner. MitoTimer is coding for DyRed1-E5 protein (a mutant form of DsRed protein) which is sensitive to free radicals. It changes its fluorescence colour from green to red in response to an increased ROS level and high mitochondrial turnover (Hernandez et al., 2013; Laker et al., 2014). Generation of the hiPSC line was performed using the Gateway® Cloning system to generate the donor vector with MitoTimer integrated between the 5′ and -3′ terminal repeats. As a second step, the piggyBac transposon system was used to drive the genomic integration of MitoTimer in chromosomal DNA for stable expression (see Figure S2).

**Generation of ACTN2-eGFP⁺ hiPSC line**

To monitor CM sarcomere contraction and relaxation in real-time we generated a α—cardiac actinin (ACTN2)-enhanced green fluorescent protein (eGFP⁺)-hiPSC line by the CRISPR-Cas9 (Sachinidis, 2021) and the homology-directed recombination (HDR) approach as we described more recently (Acharya et al., 2022). A brief description is indicated in Figure S3. The contractile velocity of c-CMs and SMG-CMs was determined by analysing the videos using the VA1.9 software (Acharya et al., 2022).
QUANTIFICATION AND STATISTICAL ANALYSIS
For all experiments, the statistical data are represented as mean ± standard deviation (SD). To calculate the p value of significance, a two-tailed Student’s t-test (t-test) from Microsoft Excel software was used and p values <0.05 were considered as statistically significant. The RNAseq data were analysed by the one-way ANOVA followed by Tukey post hoc tests for statistics using the RStudio (https://www.rstudio.com/products/rstudio/).