# Targeted biallelic integration of an inducible Caspase 9 suicide gene for safer iPSC-based cellular therapies

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#### Abstract

The teratoma forming potential of pluripotent stem cells (PSCs), and genetic aberrations that may lead to tumor formation from PSC derivatives, are considered as a major safety risk for cellular therapies. Introduction of inducible suicide genes as synthetic fail-safe systems has been proposed to minimize these risks. Recent research challenged the usefulness of such systems even for targeted introduction via accurate gene editing approaches. Apparently transgene silencing and elimination of a HTK suicide gene through "loss-of-heterozygosity" (LoH) led to cell clones that escaped the induced suicide.

We have introduced an inducible Caspase9 (iCasp-9) suicide gene into induced pluripotent stem cells (iPSCs), that has already been applied clinically in other settings. The iCASP9 gene coupled to a red nuclear GFP variant under control of the CAG promoter was inserted into the AAVS1 locus, either monoallelic or homozygous on both alleles.

Efficient induction of apoptosis *in vitro* could be induced via treatment of iCASP9 iPSCs with two chemical inducers of dimerization (CID) at different concentrations for 24 hours. While NOD*SCID* mice after transplantation of undifferentiated monoallelic iCASP9 iPSCs under the kidney capsule developed teratomas, CID treatment for three days led to rapid shrinking of such tumor structures. In individual mice, however, that received monoallelic iCASP9 iPSCs, tumor-like human tissue could be detected after CID treatment.

Further *in vitro* experiments confirmed that in very rare subclones monoallelic iCASP9 hiPSCs lost transgene expression and can became resistant to CID induction *in vitro* with frequencies of ~ 3x10<sup>-8</sup>. Analysis of CID-resistant subclones identified either elimination of the transgene, presumably via LoH, or via methylation of the CAG promoter as underlying mechanism.

In contrast, we never observed any CID resistant escapees form biallelic iCASP9 iPSC clones, even after treatment of up to 0,5x10<sup>9</sup> iPSCs. This observation further argues for LoH as underlying mechanism of transgene elimination in monoallelic clones and suggests that CAG promoter methylation on both alleles represent independent events.

In conclusion, biallelic integration of an iCASP9 safety switch in the AAVS1 locus allows for efficient induction of cellular suicide and may substantially increase the safety level of iPSC-based therapies. We propose that safety levels should be calculated by relating the observed frequencies of clonal escapees to clinically relevant cell numbers, i.e. cell number in tumors of a size that is readily detectable by modern imaging approaches.

#### Introduction

Pluripotent stem cell (PSC) technologies come out of age and a number of clinical trials applying embryonic stem cell (ESC) or induced pluripotent stem cell (iPSC)-based cell products are ongoing or in preparation. The tumorigenic potential of PSC-derived cell products, however, either in terms of contaminating undifferentiated cells that form teratoma or due to malignant transformation caused by mutations acquired and enriched during reprogramming <sup>1</sup> or culture expansion <sup>2</sup> is considered as a major safety concern.

While improved protocols for targeted differentiation of PSCs into the therapeutic derivatives of interest substantially reduced the risk for teratoma formation, introduction of fail-safe systems with drug-inducible suicide genes have been proposed to further reduce tumour risks. Such fail-safe systems include the inducible expression of the herpes simplex virus-thymidine kinase (HSV-TK), which has been clinically applied already more than 25 years ago <sup>3</sup>, and the inducible Caspase9 (iCASP9) safety switch system <sup>4</sup> that has been clinically applied more recently in T cells using a retroviral approach <sup>5</sup>.

In a first study that attempted to quantitatively define the safety level of PSC transplants with integrated fail-safe systems , mouse and human ESCs carrying an HSV-TK system controlled by one allele of the endogenous CDK1 locus were applied. Despite general functionality of the HSV-TK suicide system concerning elimination of cycling ESCs, rare proliferating mouse and human cell clones were observed *in vivo* and *in vitro* with an average frequency of 6.6x10<sup>-8</sup> that became resistant to ganciclovir. Remarkably, not only transgene silencing but also loss of heterozygosity (LoH) was identified as underlying mechanism for failure of the integrated drug-inducible suicide system. Finally, the authors demonstrated that in case of ESCs with biallelic (homozygous) integration of HSV-TK in the CDK1 locus, no resistant clones appeared among a total of 1.2x10<sup>8</sup> cells pointing to a substantially higher safety level.

One potential disadvantage of the applied HSV-TK system is that HSV-TK as viral transgene may be immunologically recognized *in vivo* leading to killing of HSV-TK expressing cells even without application of ganciclovir. It should be stressed that placing a HSV-TK transgene under control of a cell cycle-dependent locus such as CDK1 (cyclin-dependent kinase 1) <sup>6</sup> indeed restricts the targeted elimination to cycling cells such as tumour cells, but is probably not applicable if the iPSC-derived graft has to proliferate in order to fulfil its therapeutic effect, because all cycling cells that express the HSV-TK gene may become target of the host's immune response. In general, the HSV-TK fail-safe system is especially effective in fast-dividing cells and may fail to eliminate populations of slowly dividing tumour cells. In addition, the use of the prodrug ganciclovir as potent antiviral drug e.g. for treatment of serious herpes virus infections may not be possible in affected patients.

In this study, we aimed to quantitatively define the safety level of hiPSC transplants with

another integrated drug-inducible suicide system, the iCASP9 safety switch. In this system, the Caspase activation and recruitment domain of human Caspase 9, a gene expressed in fetal and adult tissue that is upregulated during induction of apoptosis, has been replaced by a modified dimerizer-binding domain of the FK506-binding protein 12<sup>5</sup>. This system requires caspase dimerization after application of an otherwise bioinert small-molecule AP1903 (Chemical Inducer of Dimerization, CID) for activation of intracytoplasmic caspase-3 and induction of apoptosis <sup>4,7</sup> (**Figure 1A**). Because Caspase 9 is an endogenous gene, it can be considered non-immunogenic.

In order to ensure safe application of the safety switch and efficient cell killing also in slowly dividing iPSC derivatives we have integrated the iCASP9 construct into the AAVS1 (Adeno-Associated-Virus Site 1) safe harbour locus <sup>8</sup> under control of a synthetic promoter consisting of the cytomegalovirus early enhancer element and chicken beta-actin (CAG) promoter, known for robust expression in undiffentiated iPSCs and their differentiated progeny <sup>9</sup> (Figure 1B). In consideration of the results of Liang et al. <sup>6</sup> we have generated hiPSC clones either heterozygous or homozygous for the integrated iCASP9 safety switch. Potential emergence of cell clones escaping the induced suicide was monitored *in vivo* and *in vitro* among therapeutically relevant cell numbers.

#### Results

Generation of Human iPSC Lines with heterozygous or homozygous (mono- or biallelic) integration of an iCASP9 suicide gene coupled to a Tomato fluorescent reporter. In order to enable inducible induction of apoptosis, we applied transcription activator-like effector nucleases (TALEN)-mediated gene editing to integrate an iCASP9 safety switch into the AAVS1 safe harbor locus. iCASP9 gene was place under control of the CAG promoter. A dTomato-fluorescent protein fused to a nuclear membrane localization signal (dTomato<sub>nucmem</sub>) was utilized to facilitate indirect monitoring of iCASP9 expression and discrimination from autofluorescence, and was coupled to the iCASP9 gene via a 2A site (Figure 1B). Allelespecific PCR analysis confirmed site-specific integration into one or both alleles of the AAVS1 locus of Phoenix <sup>10</sup> and iCBPSC2 <sup>11</sup> lines (Figure S1A&B). For all further procedures and analyses, one correctly targeted monoallelic (heterozygous) and biallelic (homozygous) clone per cell line was chosen. These clones showed considerable levels of dTomatonucmem expression (Figure 1C), absence of chromosomal aberrations as demonstrated by karyotyping (Figure 1D) expressed typical pluripotency markers, differentiated into derivatives of all three germ layers and formed teratoma upon injection into NODSCID mice (Figure S2). For clarity, all selected iCASP9 cell lines that were applied thereafter are abbreviated according to their origin and genotype: monoallelic iCASP9 Phoenix, biallelic iCASP9 Phoenix, monoallelic iCASP9 hCBiPS2, biallelic iCASP9 hCBiPS2.

Chemical inducers of dimerization (CIDs) efficiently induce apoptosis in monoallelic and biallelic iCASP9 iPSC clones. Undifferentiated cells of monoallelic and biallelic iCASP9 Phoenix and iCASP9 hCBiPS2 clones cultivated as monolayer on geltrex were treated for 24h with different concentration of two CID variants AP1903 and AP20187 before staining of viable cells with Calcein staining. Determination of the proportion of Calcein<sup>pos</sup> cells by FACS revealed efficient killing of cells of all four mono- / bi-allelic cell lines with both CID variants at concentration  $\ge 0,1nM$  (Figure 2).

In vivo CID treatment of preformed teratoma after injection of monoallelic iCASP9 hiPSCs eliminates human cells and teratoma in most but not all mice. In vivo experiments in NODSCID mice were conducted to confirm the above results in vivo (Figure 3A). All mice after injections of iPSCs under the kidney capsule showed massive increase of girth as typical sign for teratoma formation after 8 weeks. Mice that received non transgenic Phoenix wildtype cells and subsequent CID treatment, as well as mice after injection of monoallelic iCASP9 Phoenix iPSCs followed by vehicle-treatment all developed teratoma that stained positive for human nuclear antigen (Figure 3B). In contrast, in three out of five mice injection of transgenic monoallelic iCASP9 Phoenix iPSCs followed by treatment with CID resulted in almost complete elimination of frequently cystic teratoma structures (Table 1 and Figure 3B). In these mice, only some abnormal puffy mouse tissue with eosinophilic infiltration around the kidney could be detected (upper middle panel), probably representing fibrotic mouse tissue that formed in response to the massive CID-induced cell death and the resulting infiltration of phagocytes and granulocytes followed by pro-fibrotic cytokine release. In the remaining two out of five mice also extensive elimination of frequently cystic teratoma structures was observed. However, in these mice also tumor like tissue was detected around the kidney that stained positive with an anti-human nucleoli antibody, suggesting that despite killing of the vast majority of engrafted cells, re-growths of escapees occurred that were apparently resistant to CID-induction of iCASP9-mediated suicide.

*In vitro* CID treatment led to selection of rare CID-resistant cell subclones from monoallelic but not biallelic iCASP9 iPSC clones. In order to further explore the frequency of the observed CID resistance in monoallelic versus biallelic iPSCs, iCASP9 hCBiPS2 and iCASP9 Phoenix cells were treated with CID (concentrations from 0,5 nM-10nM AP1903 / AP20187), seeded onto irradiated feeder cells and were cultivated for three weeks to promote the propagation of potentially surviving cells. In correspondence to the *in vivo* experiments, survival of extremely rare CID resistant cell clones was observed in several independent experiments with overall frequencies of 3.6x10<sup>-8</sup> for hCBiPSC or 2.5x10<sup>-8</sup> for Phoenix iPSCs.

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We never observed surviving clones in biallelic iCASP9 iPSCs despite of the high numbers of treated cells (Figure 4, table 2).

Rare monoallelic iPSC subclones become CID-resistant due to transgene elimination, probably via Loss of Heterozygosity (LoH), or silencing via CAG promoter methylation. Twelve CID resistant subclones derived from monoallelic iCASP9 iPSCs (Table 2) were further cultivated to investigate mechanisms of the acquired CID resistance. Three out of four subclones derived from iCASP9 Phoenix did not show any dTomato expression. PCR based analysis of the genomic sequence of the transgene cassette in the AAVS1 locus of these iCASP9 Phoenix subclones (#1, #2 and #4 in Figure 5B) revealed elimination of the complete transgene, very likely via LoH. In the remaining clone with relatively weak dTomato expression neither transgene elimination nor increased methylation of CpGs in the CAG promoter or the adjacent AAVS1 (PPP1R12C) locus (Fig S4) as analyzed by Nanopore sequencing was observed (data not shown). In this case, the underlying mechanism remained elusive.

Loss of the transgene could not be demonstrated in any of the eight analyzed iCASP9 hCBiPS2 subclones. Interestingly, repeated CID-treatment led to elimination of dTomato expressing cells in all these CID-resistant subclones, however, dTomato expressing cells re-appeared in all subclones several days after CID expression (Figure S3) suggesting reversible epigenetic mechanisms to be responsible for transgene silencing.

Exemplarily, two of these phenotypically similar subclones (#4 and #7) were analyzed for increased methylation of CpGs in the AAVS1 locus or the CAG promoter via Nanopore sequencing. The observed high methylation rate of the CAG promoter in these subclones (**Figure 5C**) perfectly correlates with loss of dTomato expression and their resistance towards CID-treatment. Again, no elevated methylation of the adjacent AAVS1 (PPP1R12C) locus was observed (data not shown).

#### Discussion

The availability of human ESCs and iPSCs with their far-reaching potential for proliferation and differentiation offers novel opportunities for the development of tailored cellular therapies. Therapeutic risks associated with PSC-based therapies include ones that are related to abnormal cell functions such as arrhythmia in case of cardiomyocytes or dysregulation of hormones in pancreatic islands or dopaminergic neurons, but especially tumor formation is a major concern. Integration of synthetic fail-safe systems into PSCs have been proposed as additional safety measure enabling induced suicide in case of transplant-associated tumor formation. Whereas in case of retro- and lentiviral delivery of such systems, the appearance of suicide-resistant

subclones is not surprising considering the frequently observed high levels of transgene silencing, targeted integration into genomic loci known for their robust expression was considered as more appropriate approach to achieve reliable induced elimination of transplanted cells. A recent study, however, challenged that estimation. With a total of eight suicide-resistant escapee clones among a total of 120 million cells for a monoallelic HSV-TK suicide gene integrated into the endogenous CDK1 locus, Liang et al. observed an overall frequency of  $6.7 \times 10^{-8}$  escapees <sup>6</sup>.

With a total of 16 escapee clones among 500 million cells (3.2x10<sup>-8</sup>) we have now observed a comparable frequency for the CAG-iCASP9 safety switch inserted in the AAVS1 locus. Similar to Liang et al., we never observed resistant cell clones that develop from iPSCs carrying a homozygous (biallelic) safety switch. In that configuration, elimination of the suicide gene via LoH should be impossible.

In case of the observed aberrant promoter methylation as underlying mechanism for safety switch inactivation the question remains whether such rare events depend on the status of the cellular methylation machinery, which would imply that in the event of methylation of one allele there is also an increased likelihood for similar methylation of the second allele. The fact, however, that despite a high number of 0.8 billion cultured iPS cells we never observed any escapee from two iPSC lines with integrated biallelic iCASP9 safety switch suggest that the observed aberrant methylation of the promoter that occurs in very rare cases ( $\sim 3x10^{-8}$ ) on one allele is completely independent of the second allele. If this hypothesis applies, a simultaneous methylation of both alleles would be extremely unlikely ( $3x10^{-8} \times 3x10^{-8} = 9x10^{-16}$ ).

In theory, also individual loss of function mutations may lead to CID resistance. Homologous recombination may even lead to transmission of the mutation to the 2<sup>nd</sup> allele. While such events are also considered very rare and were not observed in our experiments, they certainly deserve further research. Also further mechanisms such as gene silencing due to histone modifications have to be considered.

Our results indicate that even targeted integration of a monoallelic safety switch into a safe harbor site known for robust transgene expression is insufficient to exclude development of CID-resistant subclones. Although a biallelic approach most likely offers much higher safety levels, the question remains how to calculate the safety level not only for a specific cell product but also for a given clinical scenario. Addressing that question, Liang et al. developed the term *safe-cell level (SCL)* as *the number of* 

therapeutic batches in which there is expected to be one none-safe batch <sup>6</sup>. This SCL, which is actually reflecting the frequency of escapees, was then put in relation to clinically relevant cell numbers in terms of the therapeutically applied cell doses. These will often be substantially higher than typical cell numbers handled in conventional cell culture systems and may range from  $10^5$  to  $10^{10}$  cells, e.g. it is estimated that ~  $10^9$  cardiomyocytes would have to be replaced after myocardial infarction.

It is debatable, however, whether relation of a calculated SCL to the size of a required therapeutic cell dose is the most appropriate approach to estimate the safety level of a given cell product. Integrated suicide genes are mainly considered as safety measure to eliminate proliferating tumor cells that either arise as teratoma from contaminating undifferentiated pluripotent stem cells, or develop from rare cell clones that carry mutations which lead to altered expression or function of oncogenes. It is, however, extremely unlikely that a very rare cell clone within the therapeutic cell batch that acquired CID resistance either through LoH, silencing of the transgene or other randomly developed mutation, undergoes another very rare event, which is tumor transformation.

More relevant are other scenarios, where *in vivo* single contaminating undifferentiated PSCs among the therapeutic cell transplant form a teratoma, or an individual cell clone acquires (a) genetic aberration(s) and undergoes tumor transformation, in both cases followed by massive proliferation. Among the generated large number of tumor cells, clonal CID resistance may develop as second, independent and again very rare, event. We therefore consider it more reasonable to relate the frequency of suicide-resistant escapee clones to the number of cells in a tumor of a size that can be routinely detected by modern imaging approaches, than to the size of a produced cell batch or the therapeutic cell dose. Actually, recent estimations for cell numbers in tumors tissue range from  $10^7$ - $10^9$  cells for a tumor of 1 cm<sup>3</sup>, a size that should be reliably detectable by modern imaging approaches <sup>12</sup>. Large tumors may even contain more than  $10^{10}$  cells. Provided a frequency of  $\sim 5x10^{-8}$  for clonal suicide escapees in cells with a monoallelic safety switch and  $10^9$  cells in a tumor, it is obvious that it is not unlikely that a tumor when it becomes large enough to become detected clinically may already contain CID-resistant tumor cell clones.

While it appears clear that monoallelic integration of synthetic safety switches is insufficient to provide an increased safety level for cell therapy products, large mass production settings would be required to further explore the frequency of clonal

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escapees from cells with integrated biallelic safety switches. Experimental proof, however, of the hypothesized independency of suicide transgene inactivation on both alleles, which would imply extremely low frequencies of clonal escapees in a range of 10<sup>-15</sup> is almost impossible. Since even with most advanced mass culture technologies production of not more than 10<sup>7</sup> cells/ml is possible <sup>13</sup>, a culture volume of 100.000l would be required to generate the huge number of 10<sup>15</sup> cells.

#### **Material and Methods**

**Plasmid construction.** The used AAVS1 TALENs were generated via the golden gate assembly method {Morbitzer, 2011 #48} and contain wild-type FokI nuclease domains. The AAVS1 locus-specific TALEN expression cassettes <sup>14</sup> were placed under control of a CAG promoter.

The AAVS1.iCaspase9 donor was based on the vector SFG.iCasp9.2A.∆CD19 described in Stasi et al 2011. For generation of the AAVS1 donor plasmid standard cloning technologies were used. The AAVS1 targeting vector contains CAG-iCasp9-2A-dTomato<sub>nucmem</sub>-RbGpA flanked by two arms of ~700 bp AAVS1 locus homology sequences in a pUC19 expression vector backbone (Thermo Fisher Scientific).

As described in <sup>5</sup>, the transgene iCasp9 consists of the sequence of the human FK506-binding protein (FKBP12; GenBank number, AH002818) with an F36V mutation, connected through a Ser-Gly-Gly-Gly-Ser linker to the gene encoding human caspase 9 (CASP9; GenBank number, NM001229).

**Cell culture.** We used two human iPSC (hiPSC) lines generated in our group. hCBiPS2 is based on the lentiviral transduction of cord-blood-derived endothelial cells {Haase, 2009 #50} and Phoenix is based on cord-blood-derived CD34+ cells transduced with sendai virus vectors {Haase, 2017 #56}. The hiPSC lines were cultured and expanded on irradiated mouse embryonic fibroblasts (MEFs) in iPS medium (knockout Dulbecco's modified Eagle's medium supplemented with 20% knockout serum replacement (KSR), 1 mM L-glutamine, 0.1 mM b-mercaptoethanol, 1% nonessential amino acid stock (all from Thermo Fisher Scientific), and 10 ng/ml basic fibroblast growth factor (bFGF; supplied by the Institute for Technical Chemistry, Leibniz University, Hannover, Germany) {Chen, 2012 #51}).

**Transfection and clone establishment.** For Nucleofection hiPSCs were expanded as monolayer cultures on Geltrex (Thermo Fisher Scientific), cultivated in mTESR1 (STEMCELL Technologies) and harvested by Accutase (Thermo Fisher Scientific). The transfection was performed with Neon® Transfection system (ThermoFisher Scientific). 1×10<sup>6</sup> cells were resuspended in 105 µL Neon® buffer, electroporated with 5 µg of each plasmid encoding for AAVS1- specific-TALEN and 15 µg donor plasmid with two pulses at 1,000 V for 20 ms, and plated onto Geltrex-coated dishes with MEF conditioned medium (CM) (DMEM/F12 supplemented with 15% KSR, 100µM β-mercaptoethanol, 1% nonessential amino acid stock (all from Thermo Fisher Scientific), 10ng/mL bFGF and 10 mM Y-27632 (both from the Institute for Technical Chemistry, Leibniz University Hannover).

Transfected cells were cultivated as a monolayer prior to Fluorescence Activated Cell Sorting (FACS). For clone generation, cells were harvested from the monolayer culture by Accutase

on day 10 after transfection, and were sorted on the FACSAria IIu (BD Bioscience) or XDP (Beckman-Coulter) for of dTomato<sub>nucmem</sub> positive cells. Sorted populations were plated at low density onto Geltrex-coated dishes. Colonies were picked manually and transferred into feeder-based culture conditions resulted in single-cell clones, which were also analyzed via PCR screening (see schematic illustration in Figure S1).

**PCR Screening and characterization of transgenic clones.** Genomic DNA was prepared using the QIAamp DNA Blood Mini Kit (QIAGEN) according to the manufacturer's instructions, and 100ng of gDNA was amplified by PCR with GoTaq DNA polymerase (Promega). Sequences and specifications of the primers are shown in Table S1.

**Karyotype analysis.** After treatment of adherent hiPSC for 30 min with colcemid (Invitrogen), cells were trypsinised and metaphases were prepared according to standard procedures. Fluorescence R-banding using chromomycin A3 and methyl green was performed as previously described in detail (Schlegelberger et al., 1999). At least 15 metaphases were analyzed per clone. Karyotypes were described according to the International System for Human Cytogenetic Nomenclature (ISCN).

**Immunohistochemistry for characterisation of transgenic clones.** Cells were fixed with 4% paraformaldehyde (w/v), permeabilized with triton blocking solution and stained by standard protocols using primary antibodies, listed in Table S2, and appropriate secondary antibodies. Incubation with primary antibodies and corresponding isotype controls was performed overnight at 4 °C. Staining of living cells for extracellular markers was performed for 1 h. Secondary antibody staining was performed afterwards. Cells were counterstained with DAPI (Sigma) for the analyzes with an AxioObserver A1 fluorescence microscope and Axiovision software 4.71 (Zeiss) or Propidium Iodid (PI) (final concentration 1µg/mI) (Thermo Fisher Scientific) for the analyses using a MACSQuant® Analyzer (Miltenyi Biotech). Flow cytometric data evaluation was done with FlowJo 7.6.5 software (Celeza).

*In vitro* differentiation. HiPSC cultured on MEFs in iPS medium were detached with CollagenaseIV (Thermo Fisher Scientific), dispersed in small clumps and cultured in differentiation medium (80% IMDM supplemented with 20% fetal calf serum, 1mM L-glutamine, 0,1mM ß-mercapthoethanol and 1% nonessential amino acid stock) in ultra-low-attachment plates (Corning Inc., NY, USA) for 7 days. On day 7 the formed embryoid bodys (EB) were plated onto 0,1% gelatin coated tissue culture dishes and cultured for a further 13 days before fixation and immunostaining.

#### Induction of apoptosis with the Chemical Inducer of Dimerization AP20187 and AP1903.

iPSCs were seeded on Geltrex- coated tissue culture plates, cultivated in mTESR1 and grown 48hrs before functionality of the suicide gene was assessed by adding the chemical inducer of dimerization (CID) (AP20187, B/B Homodimerizer, Clontech Laboratories, Inc), or AP1903. AP1903 was synthesized in house at the Institute for Technical Chemistry, Leibniz University, Hannover, Germany. CIDs were applied for 24 h with concentrations of 0,001nM, 0,1nM, 1nM, and 10nM. For flow cytometric analysis, iCASP9\_hiPSCs not treated with CID or the hiPSC wildtype were dissociated into single cells using Accutase. Samples based on iCASP9\_hiPSC treated with CID in different concentrations were already in suspension.

For detection of living cells undifferentiated hiPSCs were stained with 1µmol Calcein AM (Thermo Fisher Scientific) for 45 minutes and additionally all samples were in parallel analyzed using PI. Data were acquired on MACSQuant® Analyzer and analyzed using FlowJo software (Flowjo, LLC).

**Teratoma assay.** For in vivo testing of iCASP9\_suicide switch functionality, the iCASP9\_hiPSCs were cultivated as colonies on MEFs in iPS medium. After detachment with CollagenaseIV the cell suspension was mixed in ratio 1:1 with matrigel (Thermo Fisher Scientific) to increase cell survival. Teratomas were induced by injection of the cell/matrigel suspension (~1 million cells per injection) under the renal capsule of NOD/*SCID* mice. After 8– 10 weeks, the animals showed increased abdominal girth and body weight.

Then, the mice were administrated 50µg/day/animal (2,5mg/kg) AP1903 (4% AP1903 stock solution (3,125mg/ml), 10% PEG 400 (100%), and 86% Tween (2% in water)) or the same amount of PEG400/Tween solution only as vehicle. The injection was given intraperitoneal (IP) for three days. Nine days after last injection mice were sacrificed. During all experiments, the "Principles of laboratory animal care" (NIH publication No. 86-23, revised 1985) as well as the Animal Welfare Law of Lower Saxony were followed.

**Histological Analysis and Confocal microscopy.** For histological analyses, the samples were fixed in formalin and processed for paraffin embedding according to standard protocols. Histological staining was conducted on paraffin cuts using hematoxylin and eosin (H&E).

For immunofluorescence analyses of human cells, tissue slides of paraffin were used. Antigen retrieval was achieved by preheated sodium citrate buffer according to standard protocols. After antigen retrieval, tissue slides were stained using the protocol described before for immunohistochemistry.

Confocal images were acquired with a Zeiss LSM 780 and ZEN black 2.3. To separate the immunostaining signal from autofluorescence background online fingerprinting mode was used. For that reference spectra were taken from single fluorescence staining. Several

autofluorescence reference spectra were taken from specific regions of non-stained samples and used for unmixing. Images were obtained with a 20x/0.8 Plan-Apochromat, 405 nm and 488 nm excitation at zoom 0.6 (image dimensions  $1024 \times 1024$  pixel, 700 x 700 µm).

To correctly compare immunostainings all samples were prepared in parallel and imaged at identical acquisition settings. Images were displayed at the same brightness settings to emphasize the specificity of the conditions. A faint gauss blur was applied to the images.

*In vitro* testing for CID treatment surviving subclones. IPSCs were seeded on Geltrexcoated tissue culture plates or flasks, cultivated in mTESR1 and grown until a confluent monolayer is reached. Medium was changed to medium containing the chemical inducer AP20187 or AP1903 in concentrations from 0,5nM up to 10nM. After 24 hours, the complete medium containing all treated cells were centrifuged and transferred into feeder-based culture conditions. The treated hiPSCs were cultured in iPS medium (described in Cell Culture). In the first 24 hours the medium was supplemented with 10 mM Y-27632 to allow the survival of single cells. After 3 weeks of cultivation surviving colonies were picked, cultivated further under feeder-based culture conditions. After prolonged cultivation surviving subclones were transferred to feeder-free- conditions described before and were treated again with 10nM AP1903 (see also illustration Figure 5). The achieved population was analyzed for rate of methylation via Nanopore Sequencing and as described before (PCR Screening) for correct integration of the iCASP9-Donor construct (illustration Figure 5). Sequences and specifications of the primers are shown in Table S1.

**Nanopore sequencing.** Genomic DNA samples were collected before and after the treatment with AP1903 and stored as a pellet at minus 80°C. Isolation of genomic DNA was done using the NucleoBond® HMW DNA Kit (Macherey Nagel) according to the manufacturer's instructions. For Cas-mediated PCR-free enrichment of the construct integrated between base position 55115763 and 55115769 on chromosome 19 of hCBIPSC and Phoenix IPSC guideRNAs were designed by means of the "Alt-R Custom Cas9 crRNA Design Tool" (IDT®). For this purpose, the sequences of the regions, in which the potential guide RNAs should be located in order to obtain fragments of approximately 5.000 base pairs length, were pasted into the dialogue box. The program outputs alternative guide RNAs in the inserted region and indicates the respective on-target potential (values from 0 to 100, the higher the better) and off-target risk (values from 0 to 100, the higher the lower the off-target risk). Only guides with an on-target-potential ≥ 60 and an off-target risk ≥ 75 were accepted for our sequencing experiments and were ordered from IDT (Integrated DNA Technologies, Coral-ville, USA). During establishment of the best guideRNAs should be positioned in relation to

the target (for more detail see also https://biorxiv.org/cgi/content/short/2021.09.17.460763v1), a combination of the guideRNAs no. 2134, 2136, 55 and 2139 (tableS3) proved to be most effective as it yielded the highest number of calls per CpG-side. It is of utmost importance to obtain а sufficient number of calls per CpG-site (https://biorxiv.org/cgi/content/short/2021.09.17.460763v1) in order to be able to make a valid statement about the methylation rates of the CpGs. The respective sequences of the guides and their position in the construct are shown in figure S4. To enable sequencing of this CGrich and therefore difficult to sequence area, the DNA had to be cut into shorter pieces of about 5.000 base pairs each in order to avoid the formation of secondary structures. Before sequencing, the DNA quality was assessed in order to ensure that only high-molecular weight DNA was used in the subsequent library preparation for the Nanopore run. For this purpose, a pulsed-field gel analysis was carried out using a Pippin Pulse electrophoresis power supply (Sage Science, Beverly, USA). 5 µg of DNA were used per sample for the library preparation by the Cas-mediated PCR-free enrichment protocol. The latter includes the usage of the Ligation Sequencing (SQK-LSK109) Kit (Oxford Nanopore Technologies, Oxford, UK) and the Native Barcoding Expansion 1-12 (PCR-free) (EXP-NBD104) Kit (Oxford Na-nopore Technologies, Oxford, UK). The entire process consists of the following steps: Dephosphorylation of genomic DNA, Preparation of the Cas9 ribonucleoprotein complexes (RNPs), Cleavage and dA-tailing of target DNA, Native barcode ligation with appropriate purification by beads and final pooling of samples, Adapter ligation (due to barcoding, AMII has been used instead of AMX) and purification by AMPure XP beads (0.3 x volume). The pellet was resuspended in 14 µl of preheated elution buffer at 37°C for 20 minutes and 12 µl of the eluat were added to sequencing buffer and loading beads for the final library preparation step. Afterwards, the SpotON flow cell FLO-MIN106D (Oxford Nanopore Technologies, Oxford, UK) was primed and loaded, and the sequencing run was started on a MinION- device (Oxford Na-nopore Technologies, Oxford, UK).

For basecalling and demultiplexing data guppy basecaller v5.0.11 (Oxford Nanopore Technologies, Oford, UK) was used with standard settings and config file 'dna\_r9.4.1\_450bps\_hac'. The basecalled reads were aligned to the reference genome using minimap2 v2.20 (<u>https://doi.org/10.1093/bioinformatics/btv191</u>). For methylation calling we used nanopolish v0.13.2 (DOI: 10.1038/nmeth.4184 ).

**PCR-based analysis of Loss of Heterozygosity (LoH).** Genomic DNA was prepared using the QIAamp DNA Blood Mini Kit (QIAGEN) according to the manufacturer's instructions, and 100ng of gDNA was amplified by PCR with GoTaq DNA polymerase (Promega). Sequences and specifications of the primers are shown in Table S1.

**Statistical analyses.** Data are presented as mean  $\pm$  SD (n = 21-27) and were tested for normal distribution using theD'Agostino & Pearson omnibus normality test. As data points were not normally distributed, group comparison was performed by means of the non-parametric Kruskal-Wallis test (p<0,01) for independent samples.

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### **Author Contributions**

S.W. designed, performed and analysed experiments. S.M. and A.H. was involved in the cloning procedure of the donor. and design of guide RNAs. K.J. and M.D. designed, performed and analysed Nanopore Sequencing. S.G. analysed the H&E stainings and provided expertise and feedback. W.K. and A.M. performed and analysed the *in vivo* experiments (teratoma assay). A.K. provided Y-27632 and AP1903. A.Z. and E.P. were involved in analyses of remaining human cells and edit the resulting pictures (confocal microscopy). G.G. performed and analysed the karyotype analysis. M.A. provided expertise and feedback. N.L. helped with data interpretation. T.M. contributed to experimental design. R.Z. contributed expertise and helpul discussion. U. M. and S.W. wrote the manuscript.

### **Competing Financial Interests Statement**

The authors have no commercial, proprietary, or financial interest in the products or companies described in this article.

## **Figures and Figure Legends**

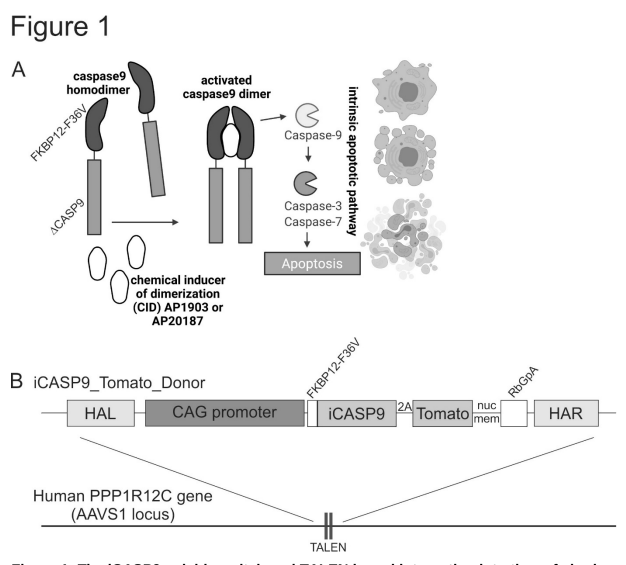
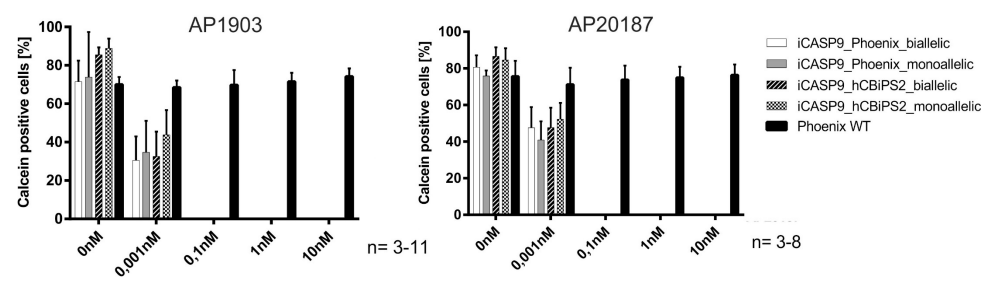


Figure 1: The iCASP9 suicide switch and TALEN based integration into the safe harbor locus AAVS1 in human iPSCs. A) Function of the iCASP9 suicide system. The CASP9 gene is deleted for its endogenous caspase activation and recruitment domain ( $\Delta$ CASP9) and was coupled to the sequence of a mutated FK506-binding protein, FKBP12, with an F36V mutation. FKBP12-F36V binds with high affinity to otherwise bioinert small-molecule dimerizing agents (chemical inducer of dimerization (CID), AP1903 and AP20187, which are lipid-permeable tacrolimus analogues. In the presence of the drugs, the iCASP9 pro-molecule dimerizes and activates the intrinsic apoptotic pathway, leading to cell death via apoptosis. (Scheme adapted from Gargett and Brown, Pharmacology 2014 p.1 Article 235).

B) Schematic illustration of iCASP9 donor construct and the AAVS1 target site. The iCASP9 donor construct consists of HAL and HAR left and right handed homology arms, a CAG promoter, an iCASP9 gene coupled via a 2A site to a dTomato fluorescence marker with a nuclear membrane localization signal. Integration of the iCASP9 donor construct was achieved via TALEN based gene editing into the AAVS1 located in the PPP1R12C gene on human chromosome 19.

Figure 2



**Figure 2: Chemical inducers of dimerization (CIDs) efficiently induce apoptosis in monoallelic and biallelic iCASP9 iPSC clones.** FACS analysis of undifferentiated iCASP9 clones cultivated on geltrex with or without CID AP1903 / AP20187 treatment for 24h. As demonstrated by loss of Calcein viability staining of iCASP9 iPSCs, efficient cell killing is achieved already by application of 0.1 nM AP1903 or AP20187 (mean ± SEM; n=3-11).

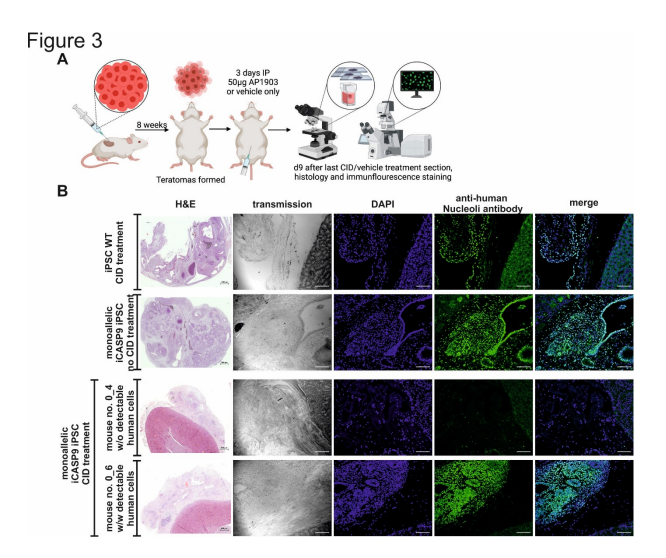
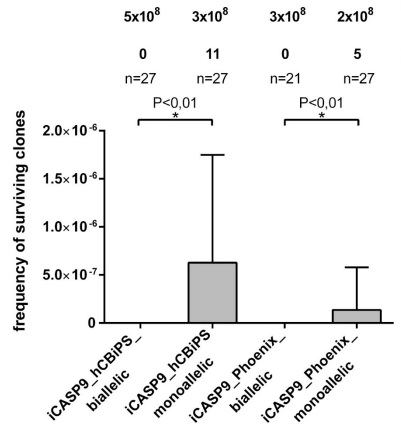


Figure 3: AP1903 eliminates preformed teratoma in the majority of NODSCID mice after injection of monoallelic iCASP9 Phoenix iPSCs but did not prevent formation of human tumor-like tissue in all animals. A) Schematic illustration of the workflow for teratoma induction and CID treatment. B) NODSCID mice were killed and sectioned 9 days after CID/vehicle treatment subsequent to injection of undifferentiated Phoenix iPSCs under the kidney capsule. (Immuno-)histology demonstrated growth of teratoma in all mice that received WT iPSCs and in monoallelic iCASP9 iPSCs without CID treatment (top and upper middle panels). In 3/5 mice that received iCASP9 iPSCs only some abnormal puffy mouse tissue with eosinophilic infiltration around the kidney could be detected (upper middle panel), probably representing fibrotic mouse tissue that formed in response to the massive CID-induced cell death and the resulting infiltration of phagocytes and granulocytes followed by pro-fibrotic cytokine release. Human tumor-like tissue was detected in 2/5 mice suggesting growth of CID-resistant cell clones (lower panel, see also table 1). Abbreviations: intraperitoneal, IP. (scale bars: 500µm for H&E, 100µm for other images)

## Figure 4



total number of CID treated cells total number of CID-resistant subclones

Figure 4: CID treatment of mono- and bialleic iPSCs led to selection of rare CID-resistant cell subclones from monoallelic but not biallelic iCASP9 iPSC clones. Monoallelic and biallelic iPSCs were treated with CID (concentrations from 0,5 nM-10nM AP1903 / AP20187), seeded onto irradiated feeder cells and were cultivated for three weeks to promote the propagation of potentially surviving cells. Survival of apparently CID resistant extremely rare cell clones was observed in several independent experiments  $(1x10^5 \text{ to } 0,5x10^9 \text{ cells } / \text{ experiment})$  with overall frequencies of  $3.6x10^{-8}$  (hCBiPSC) or  $2.5x10^{-8}$  (Phoenix). We never observed surviving clones in biallelic transgenic iPSCs despite of the high numbers of cell treated. Data are presented as mean  $\pm$  SD (n = 21-27). D'Agostino & Pearson omnibus normality test. Kruskal-Wallis test (p<0,01)..

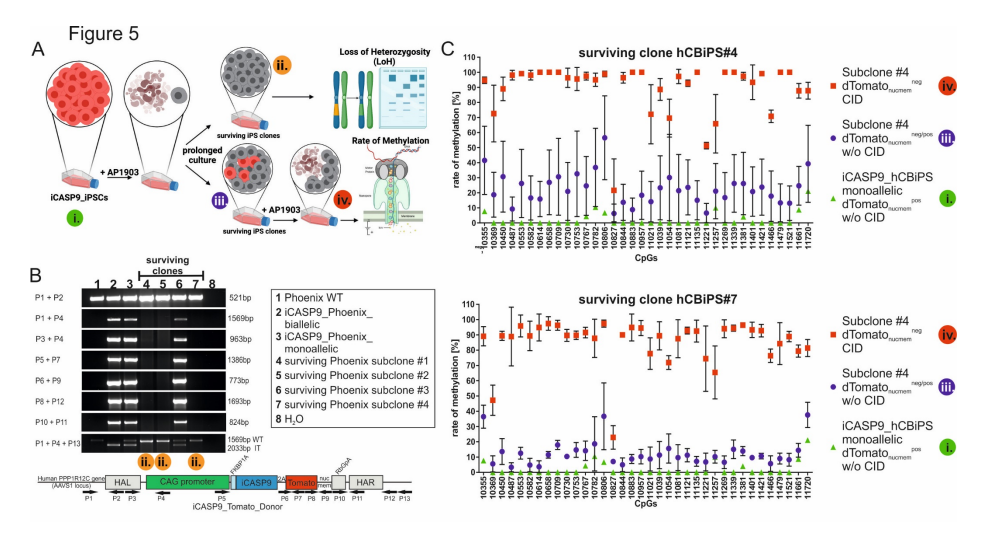


Figure 5: Rare monoallelic iPSC subclones become CID-resistant due to transgene elimination probably via Loss of Heterozygosity (LoH) or silencing via CAG promoter methylation. A) Scheme illustrating appearance and selection for Tomato<sup>neg</sup> cells resistant to CID-induced apoptosis and re-appearance of Tomato<sup>pos</sup> CID-sensitive cells during culture expansion, and analysis of different stages for LoH and methylation of

promoter elements (and surrounding genomic DNA, data not shown). Colored circles mark stages that have been further analysed for LoH and promoter methylation.

B) LoH occurred in three CID-resistant monoallelic iCASP9 Phoenix iPSC subclones but not in analyzed CID-resistant monoallelic iCASP9 hCBiPSC subclones (data not shown for hCBiPSCs). PCR based analysis of the genomic sequence of the transgene cassette in the AAVS1 locus revealed elimination of the transgene in Phoenix subclones #1, #2 and #4, very likely via LoH. Primer combinations and locations are depicted in the scheme below.

C) Nanopore Sequencing showed methylation of the CAG promoter in 2 / 2 analyzed CID-resistant monoallelic dTomato<sub>nucmem</sub> <sup>neg</sup> hcBiPSC subclones that did not show transgene elimination (data not shown). Analyses of CpG islands in the CAG promoter of surviving subclones #4 and #7 indicate strong correlation between cell survival and loss of nuclear dTomato expression after CID treatment, and high methylation rate in the CAG promoter. Data are presented as mean  $\pm$  SD (n = 3-4). A scheme of CpG islands in the AAVS1 locus and the integrated iCASP9 donor construct is shown in Figure S4. No elevated methylation of the adjacent AAVS1 (PPP1R12C) locus was observed (data not shown).

## Tables:

## Table 1: CID treatment did not reliably eliminate teratoma formed in NODSCID mice after transplantation of monoallelic iCASP9 Phoenix iPSCs.

iPSC clone	CID treatment (50µg / d / animal i.p.)	Number of mice treated*	Number of mice with detectable human cell mass after CID- treatment
monoallelic iCASP9	AP1903	5	2
monoallelic iCASP9	vehicle	3	3
wildtype	AP1903	5	5
wildtype	vehicle	3	3

\* 10<sup>6</sup> cells / mouse have been transplanted under the kidney capsule of NOD.CB17-PrkdcScid/J mice.

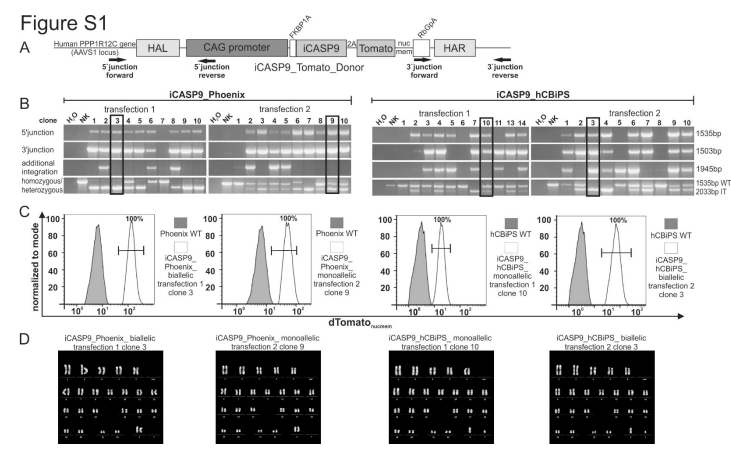
## Table 2: Rare monoallelic iCASP9 iPSC subclones become CID-resistant *in vitro* via loss-of-heterozygocity or methylation of the CAG promoter.

iPS line	Number of	Total	CID	Total	Frequency	Subclones	Subclones with	Subclones with
	CID-treated	number of	conc.	number of	of CID-	further	eliminated transgene	CAG promoter
	cells /	CID-		CID-	resistant cell	cultivated	(confirmed/tested)	methylated
	experiment	treated		resistant	subclones	and		(confirmed/tested)
		cells		subclones		analysed		
Monoallelic iCASP9 hCBiPS2	1x10 <sup>5</sup> -1x10 <sup>8</sup>	3x10 <sup>8</sup>	0,5nM-10nM	11	3,6x10 <sup>-8</sup>	8	0/8	2/2*
Monoallelic iCASP9 Phoenix	2x10 <sup>5</sup> -8x10 <sup>7</sup>	2x10 <sup>8</sup>	1nM-10nM	5	2,5x10⁻ <sup>8</sup>	4	3/4	0/1#
Biallelic iCASP9 hCBiPS2	2x10 <sup>5</sup> -5x10 <sup>8</sup>	5,2x10 <sup>8</sup>	0,5nM-10nM	0	0	0	0	0
Biallelic iCASP9 Phoenix	1x10 <sup>5</sup> -3x10 <sup>8</sup>	3,14x10 <sup>8</sup>	1nM-10nM	0	0	0	0	0

\* No further clones tested; # mechanism for CID resistance remained unclear

## Supplement

### **Supplementary Figures**



**Figure S1: Four iCASP9 iPSC clones were selected for further studies based on genotyping and flow cytometric analysis of dTomato**<sub>nucmem</sub> **expression and karyotype analysis.** A) Schematic illustration of Primer combinations and locations used for genotyping of targeted iPSCs. B) Correctly targeted and dTomato<sub>nucmem</sub> <sup>pos</sup> clones (marked with frames) were preselected based on demonstration of mono- / biallelic integration by junction PCRs on genomic DNA using primer pairs spanning the 5′- and the 3′- junction of the donor cassette and genomic AAVS1 sequence. Expression of dTomato<sub>nucmem</sub> was analyzed via flow cytometry. Histograms for dTomato<sub>nucmem</sub> expression of transgenic clones are shown exemplarily.

Karyotype analysis demonstrated absence of larger genomic aberrations. Representative metaphases are shown. Chromosome analysis was performed following standard cytogenetic procedures. At least 20 metaphases per clone were analysed.

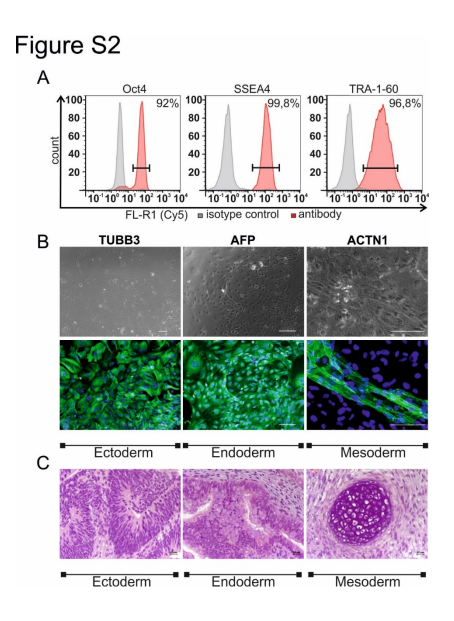
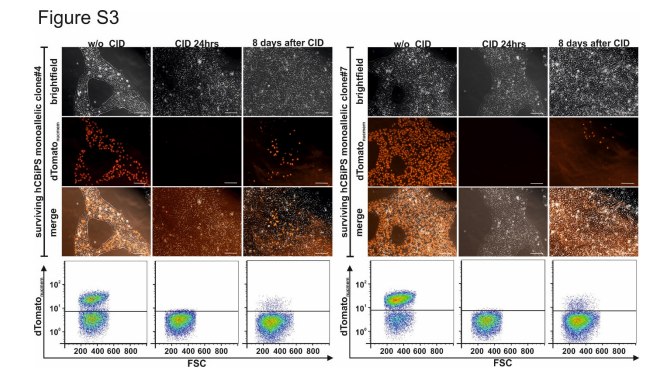


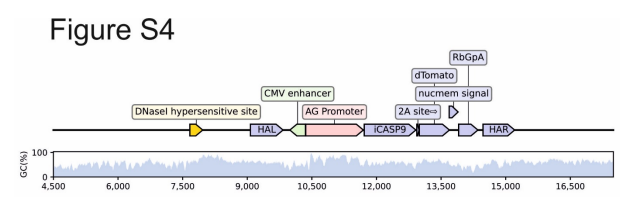
Figure S2: Characterization of iCASP9\_hiPSCs (exemplarily shown for iCASP Phoenix monoallelic). A) Immunofluorescence staining of iCASP9 hiPSCs against OCT4, SSEA-4 and TRA-1-60 and analyzes via flow cytometry demonstrate the expression of these typical pluripotency markers. B) Immunostaining of iCASP9\_hiPSCs derivatives on d21 of differentiation revealed expression of ectodermal (TUBB3), endodermal (AFP), and mesodermal (ACTN1) marker proteins (green). Nuclei are stained with DAPI (blue). Scale bars: 100 µm. C) Injection of undifferentiated iCASP9\_Phoenix\_monoallelic iPSCs into immunodeficient NODSCID mice led to formation of teratomas containing derivatives of all three germ layers. Neural tube formation representing ectodermal differentiation. Endodermal epithelium with prominent mucus-producing cells representing endoderm and mesoderm formation. Chondrocytes showing mesoderm formation. (Scale bars represent 20 mm as depicted.)



## Fig. S3: dTomato<sub>nucmem</sub><sup>neg</sup> colonies that survived CID treatment partially regain dTomato<sub>nucmem</sub> expression after prolonged cultivation.

Representative microscopic pictures and flow cytometrical analyses of two monoallelic iCASP9 hcBiPS2 subclones treated with CID are shown. Left panel: monoallelic iCASP9 subclone #4, right panel: monoallelic iCASP9 subclone #7.

Left columns: show monoallelic hcBiPS single cell clones that survived initial treatment with CID during recovery under feeder-based culture conditions. These cultures contain dTomato<sub>nucmem</sub><sup>pos</sup> and dTomato<sub>nucmem</sub><sup>neg</sup> cells (Stage iii in Figure 5A). Middle colums show the hcBiPSC clones, treated for 24 hours with CID to eliminate the dTomato<sub>nucmem</sub><sup>pos</sup> fraction. Surviving cells are dTomato<sub>nucmem</sub><sup>neg</sup> (Stage iv in Figure 5A). Starting eight days after CID-treatment new dTomato<sub>nucmem</sub><sup>pos</sup> cells could be observed (scale bar 100µm).



**Fig. S4: Schematic illustration of CpG contents in the iCASP9\_Donor integrated into the AAVS1 locus.** Content of CpGs is shown for the iCASP9\_Donor and surrounding areas of the human PPP1R12C gene.

## Supplementary Tables

Table S1: Primers and expected sizes of PCR products. Abbreviations: IT, Integrated transgene; RAS: Reverse antisense; WT, Wild type.

	Forward primer	RAS primer	Product size
5'junction	Forward CCA GCT CCC ATA GCT CAG TCT G	RAS ATG GGG AGA GTG AAG CAG AA	1535bp
3'junction	Forward CCC CTG CTG TCC ATT CCT TA	RAS GTG AGT TTG CCA AGC AGT CA	1503bp
homo-vs. heterozygous	Forward CCA GCT CCC ATA GCT CAG TCT G	RAS ATG GGG AGA GTG AAG CAG AA	1535bp WT
		RAS GTG AGT TTG CCA AGC AGT CA	2033bp IT
additional integration	Forward ATA ATA CCG CGC CAC ATA GC	RAS ATG GGG AGA GTG AAG CAG AA	1945bp
P1+P2	Forward CCA GCT CCC ATA GCT CAG TCT G	RAS GAA AAG GGA ACC CAG CGA GT	521bp
P1+P4	Forward CCA GCT CCC ATA GCT CAG TCT G	RAS ATG GGG AGA GTG AAG CAG AA	1569bp
P3+P4	Forward CTT GTA GGC CTG CAT CAT CA	RAS ATG GGG AGA GTG AAG CAG AA	963bp
P5+P7	Forward CAA CGT GCT GGT TAT TGT GC	RAS CAC CTT GAA GCG CAT GAA CT	1386bp
P6+P9	Forward GAG GAG AAT CCC GGC CCT AGG ATG GTG AGC AAG GGC GAG GAG G	RAS TTG AAT CTG TGA AGA CGG GC	773bp
P8+P12	Forward CAC CAT CGT GGA ACA GTA CG	RAS TGA AGA GCA GAG CCA GGA AC	1693bp
P10+P11	Forward CATG AAG CCC CTT GAG CATC	RAS CAG CTC AGG TTC TGG GAG AG	824bp
P1+P4+P13	Forward CCA GCT CCC ATA GCT CAG TCT G	RAS ATG GGG AGA GTG AAG CAG AA	1569bp WT
		RAS GTG AGT TTG CCA AGC AGT CA	2033bp IT

Antibody	Dilution	Company Cat # and RRID	
Mouse anti-OCT4	1:100	Santa Cruz Cat# sc-5279	
Mouse anti-TRA-1-60	1:100	Abcam Cat# 16288	
Mouse anti-SSEA-4	1:100	DSHB Cat# MC813-70 (ab16287)	
Mouse anti-α-Fetoporotein	1:300	R&D Cat# MAB1368	
Mouse anti-α-Actinin, Sarcomeric	1:800	Sigma Aldrich Cat# A7811	
Mouse anti-β-3-Tubulin	1:400	Millipore Cat #05-559	
Detection of human cells Mouse anti-human-Nucleoli		Abcam Cat# ab190710	
Cy5 Donkey anti-mouse IgG	1:200	Jackson ImmunoResearch Labs Cat# 715-175-150,	
AF488 Donkey anti-goat IgG	1:200	Jackson ImmunoResearch Labs Cat# 705-545-147	
	Mouse anti-OCT4 Mouse anti-TRA-1-60 Mouse anti-SSEA-4 Mouse anti-α-Fetoporotein Mouse anti-α-Actinin, Sarcomeric Mouse anti-β-3-Tubulin Mouse anti-human-Nucleoli Cy5 Donkey anti-mouse IgG	Mouse anti-OCT41:100Mouse anti-TRA-1-601:100Mouse anti-SSEA-41:100Mouse anti-α-Fetoporotein1:300Mouse anti-α-Actinin, Sarcomeric1:800Mouse anti-β-3-Tubulin1:400Mouse anti-human-Nucleoli1:400Cy5 Donkey anti-mouse IgG1:200	

#### Table S2: Antibodies used for immunocytochemistry / flow-cytometry

Table S3: Overview of the GuideRNAs used for the Cas-mediated PCR-free enrichment of the construct sequence for Nanopore sequencing, including the respective sequences, PAMs (Protospacer Adjacent Motifs), strand information and position in the construct.

Guide No.	Sequence	PAM	Strand	Position in the construct
2134	AGTTGGGCGCGGGGATCCGTA	AGG	+	6.578-6.597
2136	ACAATCTTCTCGACCGACAC	AGG	-	12.439-12.458

55	TTTGAGGACCTTCGACCAGC	TGG	+	12.632-12.652
2139	AATCCTACCTAACGCACTCC	TGG	-	18.076-18.095