# **Certificate of Analysis**



CELL LINE NAME	BIHi293-A	hPSCreg	Link: https://hpscreg.eu/cell-line/BIHi293-A
DONOR GENDER/AGE:	□ Male 🛛 Female □ unknown Age: 40-44		
TYPE OF DISEASE / GENETIC MODIFICATIONS	Breast Cancer		
BANK	Master Bank, MB02, Passage 18, Freezing Date 26.09.2023		
FREEZING METHOD	Bambanker		
CULTURE PLATFORM	Feeder Independent		
	Medium: E8		Coating: Geltrex
REPROGRAMMING	Sendai virus Vector details (e.g. Kit, Pub, AddgeneNr): CytoTune iPS 2.0		

TEST DESCRIPTION	Test Method	Test Specification	Result
STERILITY (viral pathogens)	<ul> <li>☑ donor tested</li> <li>□ primary cells tested</li> <li>□ iPS clone tested</li> </ul>	HBV, HCV, HIV negative	Pass
REPROGRAMMING VECTOR CLEARENCE	<ul> <li>□ parental cells tested</li> <li>□ antibody staining</li> <li>☑ PCR</li> </ul>	Vector not present	Pass
KARYOTYPE	CNV using SNP arrays	Result matches QC criteria	Pass
	G-Banding	Result matches expected karyotype	Pass
IDENTITY	STR Analysis	Identical to cells of origin	Pass
VIABILITY	Images of cells immediately post-thaw, at 48 hrs and at confluence	Growth to confluency typical of hPSCs	Pass
MORPHOLOGY	Light microscopy of cells	Typical morphology of undifferentiated hPSCs	Pass
STERILITY (mycoplasma)	Minerva Venor®GeM qOneStep	No contamination detected	Pass
STERILITY (bacteria/ yeast/ fungi)	Culture for 7 days in antibiotic free medium	No contamination detected	Pass
UNDIFFERENTIATED PHENOTYPE	Markers for undifferentiated hPSCs ⊠ IF-Staining ⊠FACS	Expression of at least three pluripotency markers detected	Pass
PLURIPOTENT DIFFERENTIATION POTENTIAL	directed differentiation	Successful differentiation to cells of all three germ layers	Pass
CONFIRMATION OF DISEASE GENOTYPE / EDITING	Sequencing of mutated site	Sequencing shows mutation	not aplicable

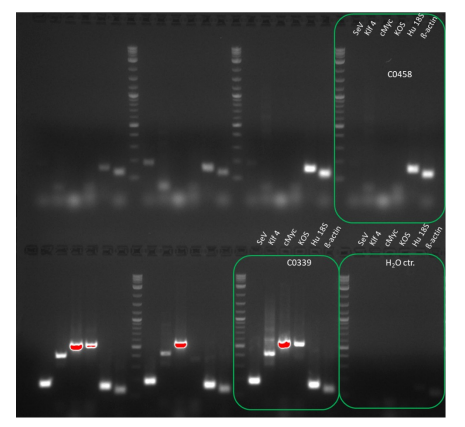
Date 29.12.23023



Cell line name	BIHi293-A
Bank ID	Clone status
Passage No.	8
Date of testing	21.07.2023
Protocol	8.4. Testing for remaining Sendai virus_CytoTune 2.0

# <u>Results</u>

2 % standard agarose gel with DNA stain Ethidiumbromid 7 $\mu L/400~mL$ 



cDNAs	Clone name/ passage		
sample			
C0458	BIHi293-A p8		
C0339	SenV pos.ctr.		
		1	
SeV	OL0452/453	181 bp	
SeV_Klf 4	OL0454/455	410 bp	
SeV_cMyc	OL00456/457	532 bp	
SeV_KOS	OL0458/459	528 bp	
Hu18sRNA	OL0107/108	152 bp	
ß-actin	OL0312/313	128 bp	

# PCR Results - Conclusion

The cell line is tested negative for Sendai virus.

Date 01.08.2023



	Reference	9		Cell line		
Sample (cell type, ID)	PBMC	42-19	9B-V4	iPSC	BIHi293-A	
Passage No.				18		
Bank ID				MB02		
DNA sample ID	D0492			D0704		
Chip-ID and Position	20621697	0037, R02	2C01	20782351	0054, R09C01	L
Date of testing	17.06.2022		27.10.202	3		
Gender (provided/estimated from chip data)	female	female	√	female	female	√

Technology:	Illumina BeadArray
Product:	Illumina Infinium Global Screening Array-24 BeadChip
Manifest:	GSAMD-24v3-0-EA_20034606_A1
Clusterfile:	GSA-24v3-0_A1_ClusterFile

#### **CNV** Analysis

Algorithm:	<b>CNV-Partition</b>
Version:	3.2.0

Parameters are set to detect copy number variations (CNVs)  $\geq$  45 kb and loss of heterozygosity (LOH) regions > 1 Mb with a confidence value > 35. Balanced translocations and inversions cannot be detected with this method. Aberrant copy number regions are identified by log R ratio and B allele frequency. Copy number changes (gains and losses) greater than **0.4 Mb** and regions of LOH above **5 Mb** are considered reportable and taken into account for interpretation. Genomic positions are based on genome build GRCh37/hg19.

If in the tested cell line (compared to the reference) new CNVs greater than **2 Mb** and/or LOH greater than **5 Mb** are detected the CNV QC test has "failed" regarding the internal QC criteria of CUSCO. We recommend not to use a "failed" cell line for further research or only after careful consideration.



	BIHi293-A 🕴	Reference (42-19B-V4)
call_rate	0.991	0.984
computed_gender	F	F
SNPs_post_filter	74.21 %	71.77 %
SNP.distance.to.ref	2	-
loss.gain_log2ratio	2.46	1.81
total_calls_CNV	13	18
total_calls_LOH	12	6
reportable_new_calls_CNV	0	0
reportable_new_calls_LOH	0	1
critical_new_calls_CNV	0	0
critical_new_calls_LOH	0	0

#### Interpretation:

The CNV analysis result suggests that the iPSC line contains neither CNVs > 2 Mb nor regions of LOH > 5 Mb.

More information can be found in the attached html report.

Further information about genes in the detected regions and linked known diseases may be provided by the UCSC Genome Browser (<u>https://genome.ucsc.edu</u>) and Decipher (<u>https://decipher.sanger.ac.uk/search</u>).

#### **References:**

1. LaFramboise, T. (1 July 2009). "Single nucleotide polymorphism arrays: a decade of biological, computational and technological advances". Nucleic Acids Research. 37 (13): 4181-4193.

2. Arsham, M. S., Barch, M. J., & Lawce, H. J. (Eds.) (2017). The AGT Cytogenetics Laboratory Manual (4th Ed.). Hoboken, NJ: John Wiley & Sons, Inc.

3. Haraksingh RR, Abyzov A, Urban AE. Comprehensive performance comparison of high-resolution array platforms for genome-wide Copy Number Variation (CNV) analysis in humans. BMC Genomics. 2017 Apr 24;18(1):321. doi: 10.1186/s12864-017-3658-x.

4. Wicell: https://www.wicell.org/home/characterization/cytogenetics/snp-microarray/single-nucleotide-polymorphism-snp-microarray-.cmsx



# G-Banding - Karyotype

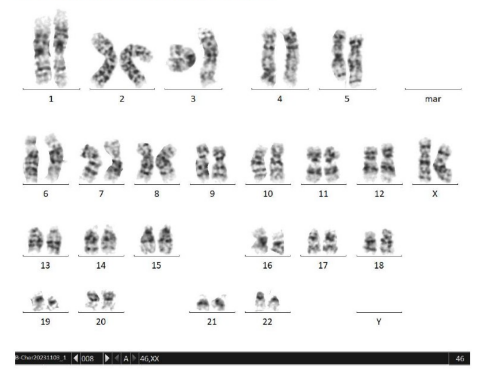
Cell line name	BIHi293-A
Bank ID	MB02
Passage No.	20
Date of testing	20.10.2023
Protocol	7.7 G-banded karyotyping

The sample preparation was carried out at BIH Stem Cell Core Facility and sent for G-bandedkaryotyping to the "Institut für Humangenetik, Universitätsklinikum Jena".

**General comments:** Karyotyping is performed using GTG stained metaphase chromosomes. With an average resolution of at least 200 bands per haploid chromosome set. Sub-microscopic changes (microdeletions/duplications) and changes <10Mb cannot be excluded by this method. Mosaics in the form of clonal changes are reported when the same change or chromosome gain occurs more than twice, and chromosome losses occur more than 3 times. A composite karyotype (cp) from 12 metaphase plates in the currently valid ISCN nomenclature is reported and a representative karyogram is provided.

#### <u>Results</u>

#### BIHi293-A p20 MB02 GBK190, Karyotyp 46,XX[cp12]



#### **Conclusion:**

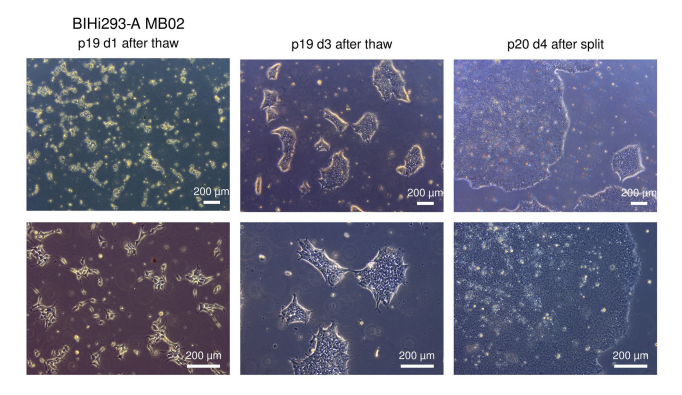
A normal female karyotype 46XX was detected for the examined sample. Only 12 metaphases could be examined.

Date: 15.11.2023

Cell line name	BIHi293-A
Bank ID	MB02
Passage No.	19
Date of testing	17.10.2023
Coating / Medium	Geltrex /E8

One vial of the cell bank was thawed and monitored during antibiotics-free cultivation. ROCK Inhibitor was used during the first 24 hours only. Cultures were evaluated regarding their morphology and viability.

#### Images:



# Conclusion:

Cells show a good post-bank recovery after thawing and form colonies exhibiting typical morphology of undifferentiated hPSCs.

Date: 24.10.2023



Cell line name	BIHi293-A
Bank ID	MB02
Passage No.	20
Test date	16.11.2023
Protocol	8.1.3 Mycoplasma testing_qPCR Minerva
Samples	<ol> <li>Negative Control (culture medium of Cell Line tested)</li> <li>Positive Control (Mycoplasma DNA from Venor® GeM qOneStep Kit)</li> <li>Cell culture supernatant from cell line</li> </ol>

#### Bacteria/Yeast/Fungi

#### Test

Cells were cultured without the addition of antibiotics over a period of 7 days. Cultures were checked daily for growth of bacteria, yeast and fungi by microscopy.

#### Results

No turbidity of the cell culture medium or microbial colonies were detected.

#### <u>Mycoplasma</u>

#### Test

Cells were cultured without the addition of antibiotics to a confluency of 80-90%. Mycoplasma contamination was tested by the qPCR-based *Venor*®*GeM qOneStep Kit*. Mycoplasma are detected at 520 nm by amplifying the 16S rRNA coding region in the mycoplasma genome. False-negative results caused by PCR inhibition are identified by the internal amplification control, detected at 560 nm.

Mycoplasma 520 nm	Internal amplification control 560 nm	Interpretation
Ct<40	Irrelevant	Sample is Mycoplasma contaminated
Ct≥40	Ct≥40	qPCR inhibition
Ct≥40	Ct<40	Sample is Mycoplasma free

#### Results

Sample	Ct of Mycoplasma DNA	Ct of Internal amplification DNA	Result
1 (neg. control)	>45	26,209	Passed
2 (pos. control)	24,375	27,433	Passed
3	>45	26,878	Negative

#### <u>Conclusion</u>

The cell line was tested negative for Mycoplasma and Bacteria/Yeast/Fungi.

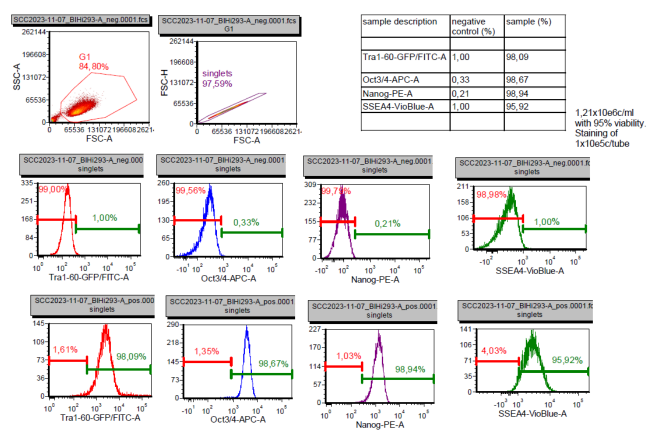
Date: 18.11.2023



Cell line name	BIHi293-A	
Bank ID	MB02	
Passage No.	24	
Date of testing	07.11.2023	
Protocol	7.14 FACS analysis of pluripotency markers	

# <u>Results</u>

20231107\_FACS analysis of markers of undifferentiated BIHi293-A MB02 p24\_one sample stained with all antibodies



# **Conclusion**

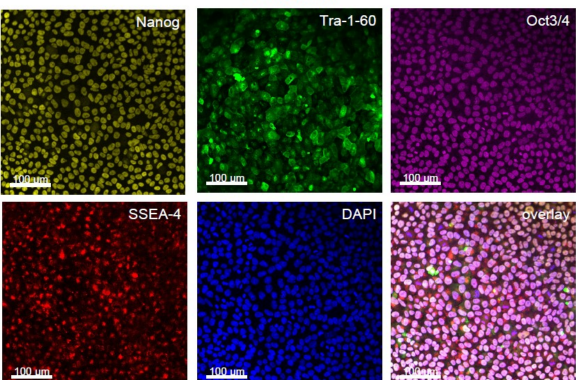
The cell line shows positive FACS results (over 80% positive) for the tested undifferentiated stem cell markers Tra1-60, OCT3/4, NANOG and SSEA-4.

Date: 07.11.2023



Cell line name	BIHi293-A	
Bank ID	MB02	
Passage No.	25	
Date of testing	20.11.2023	
Protocol	7.1 Immunofluorescence staining of markers for undifferentiated cells	

#### Results:



# BIHi293-A

# Conclusion:

The cell line shows positive staining results for the tested undifferentiated stem cell markers Nanog, OCT3/4, Tra-1-60 and SSEA4.

Date: 21/11/2023



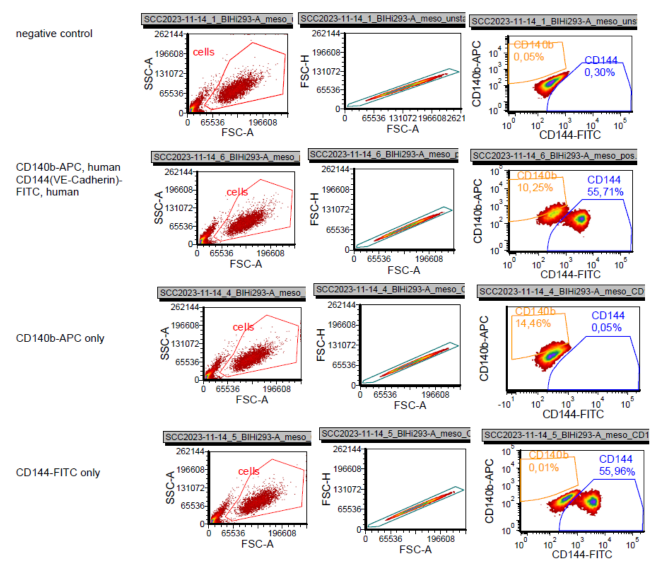
Cell line name	BIHi293-A	
Bank ID	MB02	
Passage No.	25	
Date of testing	14.11.2023	
Protocol	7.19 Validation of pluripotent differentiation potential with Trilineage	

# <u>Method</u>

Test was performed regarding the StemMACS Trilineage Differentiation Kit, human (MACS Miltenyi Biotec, Cat-No. 130-115-660). The 7-day assay enables direct differentiation of pluripotent stem cells into ecto-, meso and endoderm. The resulting cell population was measured by FACS analysis.

# <u>Result</u>

20231114 Trilineage with BIHi293-A MB02 p25 mesoderm differentiation with Miltenyi kit





# **Stem Cell Core Facility** Validation of pluripotent differentiation potential

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-10

10

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10

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10

CD184(CXCR4)-APC

196608 2621

10

131072 196608 2621

FSC-A

Gate2

65536

131072

FSC-A

98,56%

99 23%

0.54%

10<sup>2</sup> 10<sup>3</sup> 10<sup>4</sup> Anti-Sox17-Vio515

10<sup>-</sup> 10<sup>3</sup> 10<sup>4</sup> Anti-Sox17-Vio515

SCC2023-11-14\_7\_BIHi293-A\_endo\_pos.0001

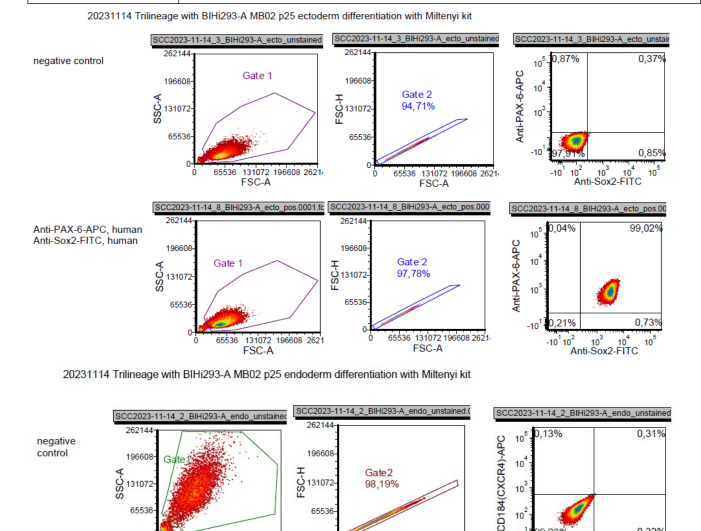
0,32%

10

89,37%

9.47%

10



65536

SCC2023-11-14\_7\_BIHi293-A\_endo\_pos.0001. SCC2023-11-14\_7\_BIHi293-A\_endo\_pos.0001.fi

FSC-H

131072 196608 2621

131072 196608 2621

FSC-A

FSC-A

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Conclusion

CD184

(CXCR4)-APC

Vio515, human

Anti-Sox17-

The cell line shows potency to differentiate into mesoderm, ectoderm and endoderm lineages. The lineage markers CD140b, CD144 (Mesoderm), Sox2, Pax6 (Ectoderm) and Sox17, CD184 (Endoderm) showed positive FACS results.

Date: 14.11.2023

65536

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