

Product Information and Testing - Amended

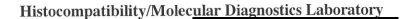
Product Information

Product Name	iPS(IMR90)-3			
Alias	iPS(IMR90) clone (#3)			
Lot Number	WB0057			
Depositor	University of Wisconsin – Laboratory of Dr. James Thomson			
Banked by	WiCell			
Thaw Recommendation	Thaw 1 vial into 4 wells of a 6 well plate			
Culture Platform	Feeder Independent			
	Medium: mTeSR™1			
	Matrix: Matrigel			
Protocol	WiCell Feeder Independent Protocol			
Passage Number	p30 These cells were cultured for 29 passages post reprogramming, at least 11 of them in mTeSR™1/Matrigel®. WiCell adds +1 to the passage number to best represent the overall passage number of cells at thaw. Fibroblasts were reprogrammed at p23.			
Date Vialed	23-September-2010			
Vial Label	WB0057 iPS(IMR90)-3 p23+30 DF 23SEPT10			
Biosafety and Use Information	Appropriate biosafety precautions should be followed when working with these cells. The end user is responsible for ensuring that the cells are handled and stored in an appropriate manner. WiCell is not responsible for damages or injuries that may result from the use of these cells. Cells distributed by WiCell are intended for research purposes only and are not intended for use in humans.			

Testing Performed by WiCell

Test Description	Test Provider	Test Method	Test Specification	Result
Post-Thaw Viable Cell Recovery	WiCell	SOP-CH-305	≥ 15 Undifferentiated Colonies, ≤ 30% Differentiation	Pass
Identity by STR	UW Molecular Diagnostics Laboratory	PowerPlex 1.2 System by Promega	Consistent with STR profile of deposited cell line	Pass
Sterility - Direct transfer method	Apptec	30744	Negative	Pass
Mycoplasma	Bionique	M250	No contamination detected	Pass
Karyotype by G-banding	WiCell	SOP-CH-003	Normal karyotype	Pass
This cell line, but not this particular	lot of material, was also tested f	or human viruses via Cha	arles River's Comprehensive Human Vir	us Panel.

Date of Lot Release	Quality Assurance Approval	
	7/14/2020	
19-January-2011	X AA	
13-3anuary-2011	AA	
	Quality Assurance	
	Signed by: Arntz, Andy	





Short Tandem Repeat Analysis*

Sample Report: 1970-STR

UW HLA#: 64095

Sample Date: 11/12/10

Received Date: 11/12/10

Requestor: WiCell Research Institute

Test Date: 11/16/10

File Name: 101116

Report Date: 11/22/10

Sample Name: (label on tube) 1970-STR

Description: WiCell Research Institute

provided genomic DNA 271.9 ug/mL; 260/280 = 1.86

Locus	Repeat #	STR Genotype
D16S539	5, 8-15	Identifying information
D7S820	6-14	has been redacted to
D13S317	7-15	protect donor
D5S818	7-15	confidentiality. If more information is
CSF1PO	6-15	required, please,
TPOX	6-13	contact WiCell's
Amelogenin	NA	Technical Support.
TH01	5-11	
vWA	11, 13-21	

Comments: Based on the 1970-STR DNA dated and received on 11/12/10 from WiCell Research Institute, this sample (UW HLA# 64095) exactly matches the STR profile of the human stem cell line iPS(IMR90) comprising 16 allelic polymorphisms across the 8 STR loci analyzed. No STR polymorphisms other than those corresponding to the human iPS(IMR90) stem cell line were detected and the concentration of DNA required to achieve an acceptable STR genotype (signal/ noise) was equivalent to that required for the standard procedure (~1 ng/amplification reaction) from human genomic DNA. These results suggest that the 1970-STR DNA sample submitted corresponds to the iPS(IMR90) stem cell line and it was not contaminated with any other human stem cells or a significant amount of mouse feeder layer cells. Sensitivity limits for detection of STR polymorphisms unique to either this or other human stem cell lines is ~5%.

Manager Date
HLA/Molecular Diagnostics Laboratory

Director Date

HLA/Molecular Diagnostics Laboratory

* Testing to assess engraftment following bone marrow transplantation was accomplished by analysis of human genetic polymorphisms at STR loci. This methodology has not yet been approved by the FDA and is for investigational use only.

File: Final STR Report

This report is confidential. No part may be used for advertising or public announcement without written permission. Results apply only to the sample(s) tested.



WiCell Research Institute

852056.A01 Page 1 of 1

December 1, 2010 P.O. #: AMENDED REPORT Original Issue Date: 11-27-10

Amendment Summary

STERILITY TEST REPORT

Sample Information:

hES Cells

1: WA15.07.07-WB0062 #1661

2: WA22-WB0046 #1491

3: WA13.C-WB0054 #7289

4: WA22-WB0053 #3855

5: iPS(IMR90)-3-WB0057 #3060

6: WA23-WB0067 #4696

7: WA15.07.03-WB0063 #8295

Date Received: Date in Test:

November 09, 2010

November 11, 2010

Date Completed:

November 25, 2010

Test Information:

Test Codes: 30744, 30744A Immersion, USP / 21 CFR 610.12 Procedure #: BS210WCR.201

TEST PARAMETERS	PRODUCT			
Approximate Volume Tested	0.5 mL	0.5 mL		
Number Tested	14	14		
Type of Media	SCD	FTM		
Media Volume	400 mL	400 mL		
Incubation Period	14 Days	14 Days		
Incubation Temperature	20 °C to 25 °C	30 °C to 35 °C		
RESULTS	14 NEGATIVE	14 NEGATIVE		

♣ A01 – Dated 12-01-10: Corrected sample information for sample # 1.

QA Reviewer

2-01-10 Date

Téchnical Reviewer

Date

Testing conducted in accordance with current Good Manufacturing Practices.





MYCOPLASM	1A TESTING SERVICES					7
APPENDIX					(2)	
Document ID#:	DCF9002F	COMMUNICATION CONTRACTOR		MARPHINIS SEES ASSESSMENT SEED AND SEED	ESTATUTE OF THE PROPERTY OF TH	AND DESCRIPTION OF THE PARTY OF
Title:	QUALITY ASSURANCE REPORT - GMP					
Effective Date:	03/12/10					
Edition #	01					

BIONIOUE® TESTING LABORATORIES, INC.

QUALITY ASSURANCE REPORT - GMP

TEST PERFORMED M-250	PROCEDURAL REFERENCE SOP's 3008, 3011, 3013	Test Performed M-700	PROCEDURAL REFERENCE SOP's 3008, 3009, 3010	
M-300 M-350 Bionique Sample ID	SOP's 3008, 3014 SOP's 3008, 3014, 3015	™-800 2785	SOP's 3008, 3011, 3016	
		02 100		

This testing procedure was performed in compliance with the FDA's Current Good Manufacturing Practice (cGMP) standards (to the extent that the regulations pertain to the procedures performed) as specified in the Code of Federal Regulations, Title 21 Parts 210 and 211 [21 CFR 210 & 211]. All related records derived from the test procedures have been reviewed by the Quality Assurance Department. The individual's signature below verifies that the methods and procedures referenced above have been followed and that the Final Report accurately reflects the raw data generated during the course of the procedures. All records, including raw data and final reports are archived on site for a minimum of seven years.

The specified test's procedures determine the intervals at which samples are inspected. The medium used for testing must pass quality control mycoplasmal growth promotion testing and sterility testing. Traceability of all of the components used is assured and supporting documentation can be supplied upon request.

Quality Assurance Review Date:		
Reviewed By QA Assistan		

NOTE:

- 1. Prior to receipt at Bionique® Testing Laboratories, Inc., the stability of the test article is the responsibility of the company submitting the sample. Bionique Testing Laboratories Inc. will assume responsibility for sample stability following receipt and prior to being placed on test.
- 2. This test is for the detection of microbiological growth and does not require statistical validation.

APPENDIX

BIONIQUE® TESTING LABORATORIES, INC.

Document ID#: DCF9002F

Title:

QUALITY ASSURANCE REPORT - GMP

Effective Date:

03/12/10

Edition#:

01

REFERENCES

Regulatory:

- Department of Health and Human Services, Food and Drug Administration (USA) [FDA]. Code of Federal Regulations [CFR], Title 21 CFR Part 210, Current Good Manufacturing Practice in Manufacturing, Processing, Packing, or Holding of Drugs; General. FDA. Office of the Federal Register, National Archives and Records Department.
- 2. Department of Health and Human Services, Food and Drug Administration (USA) [FDA]. Code of Federal Regulations [CFR], Title 21 CFR Part 211, Current Good Manufacturing Practice for Finished Pharmaceuticals. FDA. Office of the Federal Register, National Archives and Records Department.
- Department of Health and Human Services, Food and Drug Administration (USA) [FDA]. Points to Consider in the Characterization of Cell Lines Used to Produce Biologicals, Director, Center for Biologics Evaluation and Research, FDA. May, 1993. Docket No. 84N-0154.
- Department of Health and Human Services, Food and Drug Administration (USA) [FDA]. Code of Federal Regulations [CFR], Title 21 CFR Part 610.30, General Biological Products Standards; Subpart D, Test for Mycoplasma. FDA. Office of the Federal Register, National Archives and Records Department.

General:

- 1. Barile MF, Kern J. Isolation of Mycoplasma arginini from commercial bovine sera and its implication in contaminated cell cultures. Proceedings of the Society for Experimental Biology and Medicine, Volume 138, Number 2, November 1971.
- 2. Chen, T.R. In situ detection of mycoplasma contamination in cell cultures by fluorescent Hoechst 33258 stain. Experimental Cell Research, 104: 255-262, 1977.
- 3. Carolyn K. Lincoln and Daniel J. Lundin. Mycoplasma Detection and Control. U. S. Fed. for Culture Collections Newsletter, Vol. 20, Number 4, 1990.
- 4. Fetal Bovine Serum; Proposed Guideline. National Committee For Clinical Laboratory Standards (NCCLS), Vol. 10, Number 6, 1990. (NCCLS publication M25-P).
- 5. McGarrity GJ, Sarama J, Vanaman V. Cell Culture Techniques. ASM News, Vol. 51, No. 4, 1985.
- 6. Tully JG, Razin S. Methods in Mycoplasmology, Volumes I and II. Academic Press, N.Y., 1983.
- 7. Barile MF, Razin S, Tully JG, Whitcomb RF. The Mycoplasmas, Volumes 1-4. Academic Press, N.Y., 1979.
- 8. http://www.bionique.com/ Safe Cells Insights



MYCOPLASMA TESTING SERVICES

BIONIOUE TESTING LABORATORIES TNC

APPENDIX IV

DCF3013D

Document#: Edition#:

10

Effective Date:

07/15/2003

Title:

M-250 FINAL REPORT SHEET

M-250 FINAL REPORT

Direct Specimen Culture Procedure 3008, 3011, 3013

TO:

WiCell QA WiCell Research Institute

BTL SAMPLE ID#: 62784

P.O.#:

DATE REC'D:

10/12/2010

Page 1 of 2

TEST/CONTROL ARTICLE:

iPS(IMR90)-3-WB0057 #5830 p23+p31 MW

LOT#:

NA

(See Reverse)

DIRECT CULTURE SET-UP (DAY 0) 10/13/2010 DATE: INDICATOR CELL LINE (VERO) SEE DNA FLUOROCHROME RECORD SHEET DATE THIOGLYCOLLATE BROTH DAY 7 10/20/2010 DAY 28 11/10/2010 BROTH-FORTIFIED COMMERCIAL 0.5 mL SAMPLE DAY 7 10/20/2010 6.0 mL BROTH DAY 28 11/10/2010 BROTH-MODIFIED HAYFLICK DAY 7 0.5 mL SAMPLE 10/20/2010 6.0 mL BROTH DAY 28 11/10/2010 BROTH-HEART INFUSION 0.5 mL SAMPLE DAY 7 10/20/2010 mL BROTH DAY 28 6.0 11/10/2010 Document#:

DCF3013D

Edition#:

10

Effective Date:

07/15/2003

Title:

M-250 FINAL REPORT SHEET

SAMPLE ID#: 62784		AEROBIC	MICROAEROPHILIC	DATE
AGAR PLATES-FORTIFIED COMMERCIAL	DAY 7 DAY 14 DAY 21	+ 6	+ () + () + ()	$\frac{10/20/2010}{10/27/2010}$ $\frac{11/03/2010}{11/03/2010}$
AGAR PLATES-MODIFIED HAYFLICK	DAY 7 DAY 14 DAY 21	+ 8	+ 3+ + -	$\frac{10/20/2010}{10/27/2010}$ $\frac{11/03/2010}{11/03/2010}$
AGAR PLATES-HEART INFUSION	DAY 7 DAY 14 DAY 21	+ (-)	+ C+	$\frac{10/20/2010}{10/27/2010}$ $\frac{11/03/2010}{11/03/2010}$
BROTH SUBCULTURES (DAY 7) DATE: 10/20/2010				
AGAR PLATES-FORTIFIED COMMERCIAL	DAY 7 DAY 14 DAY 21	+ 🕞	+ (-) + (-) + (-)	$\frac{10/27/2010}{11/03/2010}$ $\frac{11/10/2010}{11/10/2010}$
AGAR PLATES-MODIFIED HAYFLICK	DAY 7 DAY 14 DAY 21	+ 0	+	10/27/2010 11/03/2010 11/10/2010
AGAR PLATES-HEART INFUSION	DAY 7 DAY 14 DAY 21	+ (-) + (-)	+ (-)	10/27/2010 11/03/2010 11/10/2010

RESULTS:

No detectable mycoplasmal contamination

Laboratory Director

Ph.D.

M-250 Procedural Summary: The objective of this test is to ascertain whether or not detectable mycoplasmas are present in an in vitro cell culture sample, be it a primary culture, hybridoma, master seed stock or cell line. This procedure combines an indirect DNA staining approach to detect non-cultivable mycoplasmas with a direct culture methodology utilizing three different mycoplasmal nedia formulations. The indirect approach involves the inoculation of the sample into a mycoplasma-free VERO (ATCC) indicator cell line and performing a DNA fluorochrome assay after 72-120 hours of incubation. The direct culture aspect of the test utilizes three different mycoplasmal media including both broth and agar formulations. The sample is inoculated into each of the 3 broth formulations and also onto duplicate plates (0.1 mL/plate) for each of the 3 agar formulations. Subculture from broth to fresh agar plates is carried out after 7 days incubation. Agar plates are incubated aerobically and microaerophillically in order to detect any colony forming units morphologically indicative of mycoplasmal contamination. Issuance of the final report with signature of the Laboratory Director signifies that the required controls were performed concurrently with the test sample(s) as detailed in the referenced SOPs and that all test conditions have been found to meet the required acceptance criteria for a valid test, including the appropriate results for the positive and negative controls.



BIONIQUE® TESTING LABORA	ATORIES, INC.

WIYCOPLASMA TEST	ING SERVICES
Title:	DCF3008A DNA FLUOROCHROME ASSAY RESULTS 3/24/10 07
	DNA-FLUOROCHROME ASSAY RESULTS Procedures 3008, 3009, 3011
Sample ID # 627	<u>M-250</u> Date Rec'd: <u>10/12/2010</u> P.O. #
Indicator Cells Inoc	ulated: Date/Initials: (0/14/10 / H3
Fixation:	Date/Initials: $10/18/10$ /
Staining:	Date/Initials: (0/18/10 / 13
TEST/CONTROL A	RTICLE:
iPS(IMR90)-	3-WB0057 #5830 p23+p31 MW
LOT# <u>NA</u>	
WiCell QA WiCell Resea	walk Tractitute
Wiceli Resea	Phone:
	Fax #:
ii.	
DNA FLUOROG	CHROME ASSAY RESULTS:
NEGAT	A reaction with staining limited to the nuclear region, which indicates no mycoplasmal contamination.
POSITI	YE: A significant amount of extranuclear staining which strongly suggests mycoplasmal contamination.
INCONC	CLUSIVE:
	A significant amount of extranuclear staining consistent with low - level mycoplasmal contamination or nuclear degeneration.
-	A significant amount of extranuclear staining consistent with bacterial, fungal or other microbial contaminant or viral CPE. Morphology not consistent for mycoplasmal contamination.

COMMENTS:				
Date: 19/18/10	_Results Read by:_	B	Date of Review: <u>cofreto</u>	Reviewed by: Sa



WiCell Cytogenetics Report: 003740

WISC 5830

Report Date: October 14, 2010

Case Details:

Cell Line: iPS(IMR90)-3-WB0057 (5830)

Passage #: 23+31

Date Completed: 10/14/2010
Cell Line Gender: Female

Investigator: Wisconsin International Stem Cell Bank

Specimen: iPSC on Matrigel

Date of Sample: 10/11/2010

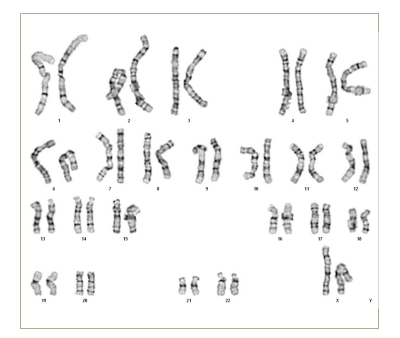
Tests, Reason for: lot release testing

Results: 46.XX

Completed by CG(ASCP), on 10/14/2010

Reviewed and interpreted by PhD, FACMG, on 10/14/2010

Interpretation: No clonal abnormalities were detected at the stated band level of resolution.



Cell: S01-01

Slide: 1-7

Slide Type: Karyotyping

of Cells Counted: 20

of Cells Karyotyped: 4

of Cells Analyzed: 8

Band Level: 475-525

Date:_____Sent To:

Results Recorded: