Thawing, Propagating, and Cryopreserving Protocol

NCI-PBCF-HTB14 (U-87 MG) Glioblastoma-Astrocytoma (ATCC®HTB-14™)

February 27, 2012; Version 1.6
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SOP: Thawing, Propagation and Cryopreservation of NCI-PBCF-HTB14 (U-87 MG)

Protocol for Thawing, Propagation and Cryopreservation of NCI-PBCF-HTB14 (U-87 MG) (ATCC®HTB-14™) glioblastoma-astrocytoma

1. Background Information on U-87 MG cell line

<table>
<thead>
<tr>
<th>Designations:</th>
<th>U-87 MG</th>
</tr>
</thead>
<tbody>
<tr>
<td>Biosafety Level:</td>
<td>1</td>
</tr>
<tr>
<td>Shipped:</td>
<td>frozen (in dry ice)</td>
</tr>
<tr>
<td>Growth Properties:</td>
<td>adherent (see appendix 1)</td>
</tr>
<tr>
<td>Organism:</td>
<td>Homo sapiens</td>
</tr>
<tr>
<td>Source:</td>
<td>Organ: brain; Disease: glioblastoma; astrocytoma; classified as grade IV as of 2007</td>
</tr>
</tbody>
</table>

For more information visit the ATCC webpage: http://www.atcc.org/ATCCAdvancedCatalogSearch/ProductDetails/tabid/452/Default.aspx?ATCCNum=HTB-14&Template=cellBiology

2. General Information for the thawing, propagating and cryopreserving of NCI-PBCF-HTB14 (U-87 MG)

| Culture Initiation | The cryoprotectant (DMSO) should be removed by centrifugation.  
|                   | The seeding density to use is about $4 \times 10^4$ viable cells/cm$^2$ or one vial of U-87 MG cells into one T-25 flask containing with 10 mL of complete growth medium EMEM + 10% (v/v) FBS. |
| Complete growth medium | The complete growth medium used to expand U-87 MG cells is EMEM + 10% (v/v) FBS  
|                       | Complete growth medium (EMEM + 10% (v/v) FBS should be pre-warmed before use by placing into a water bath set at 35 °C to 37 °C for 15 min to 30 min.  
|                       | After 30 min, the complete growth medium (EMEM + 10% (v/v) FBS) should be moved to room temperature until used. Complete growth medium (EMEM + 10% (v/v) FBS should be stored at 2 °C to 8 °C when not in use. |
| Growth Profile | Population Doubling Time (PDT) is approximately 34 h (Appendix 2). |
**Cell Growth**
- The growth temperature for U-87 MG is $37 \, ^\circ C \pm 1 \, ^\circ C$ and 5% $+ 1 \%$ CO2 in air atmosphere is recommended.

**Special Growth Requirements**
- Subculture U-87 MG cells at 75% to 85% confluence or when cell density reaches an average of $2 \times 10^5$ viable cells/cm$^2$.

**Subculture Medium**
- 0.25% (w/v) trypsin-0.53 mM EDTA (ATCC cat no. 30-2101).
- Subculturing reagents should be pre-warmed before use by placing into a water bath set at 35 °C to 37 °C for 15 min to 30 min.
- After 30 min, the subculturing medium should be moved to room temperature until used. Subculturing reagents should be stored at 2 °C to 8 °C when not in use.

**Subculture Method**
- The attached U-87 MG cells are subcultured using 0.25% (w/v) trypsin-0.53 mM EDTA (ATCC cat no. 30-2101).
- The enzymatic action of the trypsin-EDTA is stopped by adding complete growth medium to the detached cells.
- A split ratio of 1:4 to 1:5 or a seeding density of $4.0 \times 10^4$ to $5.0 \times 10^4$ viable cells/cm$^2$ is used when subculturing U-87 MG cells.

**Viable Cells/mL/Cryovial**
- The target number of viable cells/mL/cryovial is: $2 \times 10^6$ (acceptable range: $1.0 \times 10^6$ cells/mL to $2.0 \times 10^6$ cells/mL).

**Cryopreservation Medium**
- The cryopreservation medium for U-87 MG cells is complete growth medium (EMEM + 10% (v/v) FBS) containing 5% (v/v) DMSO (ATCC cat no. 4-X).

**General Procedure to be applied throughout the SOP**

**Aseptic Technique**
- Use of good aseptic technique is critical. Any materials that are contaminated, as well as any materials with which they may have come into contact, must be disposed of immediately.

**Traceability of material/reagents**
- Record the manufacturer, catalog number, lot number, date received, date expired and any other pertinent information for all materials and reagents used. Record information in the Reagent Lot Traceability Table 4 (Appendix 6).

**Expansion of cell line**
- Record the subculture and growth expansion activities, such as passage number, % confluence, % viability, cell morphology (see Figures 1, 2, Appendix 1) and population doubling levels (PDLs), in the table for Cell Expansion (Table 5, Appendix 6). Calculate PDLs using the equation in Appendix 7.

**Medium volumes**
- Medium volumes are based on the flask size as outlined in Table 1; Medium Volumes Table 1 below.

**Glossary of Terms**
- Refer to Glossary of Terms used throughout the document (see Appendix 4).

**Safety Precaution**
- Refer to Safety Precautions pertaining to the thawing, propagation and cryopreservation of U-87 MG (see Appendix 8).
### Table 1: Medium Volumes

<table>
<thead>
<tr>
<th>Flask Size</th>
<th>Medium Volume Range</th>
</tr>
</thead>
<tbody>
<tr>
<td>12.5 cm² (T-12.5)</td>
<td>3 mL to 6 mL</td>
</tr>
<tr>
<td>25 cm² (T-25)</td>
<td>5 mL to 13 mL</td>
</tr>
<tr>
<td>75 cm² (T-75)</td>
<td>10 mL to 38 mL</td>
</tr>
<tr>
<td>150 cm² (T-150)</td>
<td>30 mL to 75 mL</td>
</tr>
<tr>
<td>175 cm² (T-175)</td>
<td>35 mL to 88 mL</td>
</tr>
<tr>
<td>225 cm² (T-225)</td>
<td>45 mL to 113 mL</td>
</tr>
</tbody>
</table>

### 3. Reagents

Follow Product Information Sheet storage and/or thawing instructions. Below is a list of reagents for the propagation, subcultivation and cryopreservation of U-87 MG cells.

#### Table 2: Reagents for Expansion, Subculturing and Cryopreservation of U-87 MG cells

<table>
<thead>
<tr>
<th>Complete growth medium reagents</th>
<th>Subculturing reagents</th>
<th>Cryopreservation medium reagents</th>
</tr>
</thead>
<tbody>
<tr>
<td>Eagle’s Minimum Essential Medium (EMEM) (ATCC cat no. 30-2003)</td>
<td>Trypsin-EDTA (0.25 % (w/v) Trypsin/0.53 mM EDTA) (ATCC cat no.30-2101)</td>
<td>Eagle’s Minimum Essential Medium (EMEM) (ATCC cat no. 30-2003)</td>
</tr>
<tr>
<td>10 % (v/v) Fetal Bovine Serum (FBS) (ATCC cat no. 30-2020)</td>
<td>Dulbecco’s Phosphate Buffered Saline (DPBS); modified without calcium chloride and without magnesium chloride (ATCC cat no.30-2200)</td>
<td>10 % (v/v) FBS (ATCC cat no. 30-2020)</td>
</tr>
</tbody>
</table>

**a. Preparation of complete growth medium (EMEM + 10 % (v/v) FBS)**

The complete growth medium is prepared by aseptically combining,

1. 56 mL FBS (ATCC cat. #30-2020) to a 500 mL bottle of basal medium – EMEM (ATCC cat no. 30-2003).
2. Mix gently, by swirling.
4. Thawing and Propagation of Cells

Reagents and Material:

- Complete growth medium (EMEM + 10 % (v/v) FBS)
- Water bath
- T-25 cm² polystyrene flask
- 15 mL polypropylene conical centrifuge tubes
- Plastic pipettes (1 mL, 10 mL, 25 mL)

a. Thawing cells

Method:

1. Place complete growth medium (EMEM + 10 % (v/v) FBS) in a water bath set at 35 °C to 37 °C.

2. Label T-25 flask to be used with the (a) name of cell line, (b) passage number, (c) date and (d) initials of technician.

3. Wearing a full face shield, retrieve a vial of frozen cells from the vapor phase of the liquid nitrogen freezer.

4. Thaw the vial by gentle agitation in a water bath set at 35 °C to 37 °C. To reduce the possibility of contamination, keep the O-ring and cap out of the water. Note: Thawing should be rapid (approximately 2 min to 3 min, just long enough for most of the ice to melt).

5. Remove vial from water bath and process immediately.

6. Remove excess water from the vial by wiping with sterile gauze saturated with 70 % ethanol.

7. Transfer the vial to a BSL-2 laminar-flow hood.
b. Propagating cells

Method:

1. Add 9 mL of complete growth medium (EMEM + 10 % (v/v) FBS) to a 15-mL conical centrifuge tube.

2. Again wipe the outer surface of the vial with sterile gauze wetted with 70 % ethanol.

3. Using sterile gauze, carefully remove the cap from the vial.

4. With a 1 mL pipette transfer, slowly, the completely thawed content of the vial (1 mL cell suspension) to the 15-mL conical centrifuge tube containing 9 mL complete growth medium (EMEM + 10 % (v/v) FBS). Gently resuspend cells by pipetting up and down.

5. Centrifuge at 125 xg, at room temperature, for 8 min to 10 min.

6. Carefully aspirate (discard) the medium, leaving the pellet undisturbed.

7. Using a 10 mL pipette, add 10 mL of complete growth medium (EMEM + 10 % (v/v) FBS).

8. Resuspend pellet by gentle pipetting up and down.

9. Using a 1 mL pipette, remove 1 mL of cell suspension for cell count and viability. Cell counts are performed using either an automated counter (such as Innovatis Cedex System; Beckman-Coulter ViCell system) or a hemocytometer.

10. Record total cell count and viability. When an automated system is used, attach copies of the printed results to the record.

11. Plate cells in pre-labeled T-25 cm² flask at about 4 x 10⁴ viable cells/cm².

12. Transfer flask to a 37 °C ± 1 °C in 5 % CO₂ incubator if using flasks with vented caps (for non-vented caps stream 5 % CO₂ in the headspace of flask).

13. Observe culture daily by eye and under an inverted microscope to ensure culture is free of contamination and culture has not reached confluence. Monitor, visually, the pH of the medium daily. If the medium goes from red through orange to yellow, change the medium.

14. **Note:** In most cases, cultures at a high cell density exhaust the medium faster than those at low cell density as is evident from the change in pH. A drop in pH is usually accompanied by an increase in cell density, which is an indicator to subculture the cells. Cells may stop growing when the pH is between pH 7 to pH 6 and loose viability between pH 6.5 and pH 6.

15. If fluid renewal is needed, aseptically aspirate the complete growth medium from the flask and discard. Add an equivalent volume of fresh complete growth medium to the flask. Alternatively, perform a fluid addition by adding fresh complete growth medium to the flask without removing the existing medium. Record fluid change or fluid addition on the Cell Line Expansion Table (see Table 5 in Appendix 6).
16. If subculturing of cells is needed, continue to ‘Subculturing cells’.

- **Note: Subculture when cells are 75-85% confluent (see photomicrographs, Appendix 1).** U-87 MG cells may pile up and form clusters. They may slough from the growth surface if allowed to become over-confluent.

### c. Subculturing cells

**Reagents and Material:**

- 0.25 % (w/v) Trypsin-0.53 mM EDTA
- DPBS
- Complete growth medium (EMEM (ATCC cat no. 30-2003) + 10 % (v/v) FBS (ATCC cat no. 30-2020))
- Plastic pipettes (1 mL, 10 mL, 25 mL)
- T-75 cm², T-225 cm² polystyrene flasks

**Method:**

1. Aseptically remove medium from the flask

2. Add appropriate volumes of sterile Ca²⁺- and Mg²⁺-free DPBS to the side of the flask opposite the cells so as to avoid dislodging the cells (see Table 3).

3. Rinse the cells with DPBS (using a gently rocking motion) and discard.

4. Add appropriate volume of 0.25 % (w/v) Trypsin-0.53 mM EDTA solution to the flask (see Table 3).

5. Incubate the flask at 37 °C ± 1 °C until the cells round up. Observe cells under an inverted microscope every 5 min. When the flask is tilted, the attached cells should slide down the surface. This usually occurs after 5 min to 10 min of incubation.

   **Note: Do not leave trypsin-EDTA on the cells any longer than necessary as clumping may result.**

6. Neutralize the trypsin-EDTA/cell suspension by adding an equal volume of complete growth medium (EMEM + 10 % (v/v) FBS) to each flask. Disperse the cells by pipetting, gently, over the surface of the monolayer. Pipette the cell suspension up and down with the tip of the pipette resting on the bottom corner or edge until a single cell suspension is obtained. Care should be taken to avoid the creation of foam.

7. Using a 1 mL pipette, remove 1 mL of cell suspension for total cell count and viability.

8. Record total cell count and viability.
9. Add appropriate volume of fresh complete growth medium (EMEM + 10 % (v/v) FBS) and transfer cell suspension (for volume see Table 1 on page 4) into new pre-labeled flasks at a seeding density of $4.0 \times 10^4$ to $5.0 \times 10^4$ viable cells/cm$^2$ or a split ratio of 1:4 to 1:5.

10. Label all new flasks with the (a) name of cell line, (b) passage number, (c) date and (d) the initials of technician.

### Table 3 - Volume of Rinse Buffer and Trypsin

<table>
<thead>
<tr>
<th>Flask Type</th>
<th>Flask Size</th>
<th>DPBS Rinse Buffer</th>
<th>Trypsin-EDTA</th>
</tr>
</thead>
<tbody>
<tr>
<td>T-flask</td>
<td>12.5 cm$^2$ (T-12.5)</td>
<td>1 mL to 3 mL</td>
<td>1 mL to 2 mL</td>
</tr>
<tr>
<td></td>
<td>25 cm$^2$ (T-25)</td>
<td>1 mL to 5 mL</td>
<td>1 mL to 3 mL</td>
</tr>
<tr>
<td></td>
<td>75 cm$^2$ (T-75)</td>
<td>4 mL to 15 mL</td>
<td>2 mL to 8 mL</td>
</tr>
<tr>
<td></td>
<td>150 cm$^2$ (T-150)</td>
<td>8 mL to 30 mL</td>
<td>4 mL to 15 mL</td>
</tr>
<tr>
<td></td>
<td>175 cm$^2$ (T-175)</td>
<td>9 mL to 35 mL</td>
<td>5 mL to 20 mL</td>
</tr>
<tr>
<td></td>
<td>225 cm$^2$ (T-225)</td>
<td>10 mL to 45 mL</td>
<td>5 mL to 25 mL</td>
</tr>
</tbody>
</table>

5. Harvesting of Cells for Cryopreservation

**Reagents and Material:**

- 0.25 % (w/v) Trypsin-0.53 mM EDTA
- DPBS
- Complete growth medium (EMEM (ATCC cat no. 30-2003) + 10 % (v/v) FBS (ATCC cat no. 30-2020))
- 50 mL or 250 mL conical centrifuge tube
- Plastic pipettes (1 mL, 10 mL, 25 mL)
- Sterile DMSO
- 1 mL to 1.8 mL cryovials
- Ice bucket with ice

**Method:**

1. Label cryovials to include information on the (a) name of cell line, (b) passage number (c) date.
2. Prepare cryopreservation medium by adding DMSO to cold complete growth medium (EMEM + 10 % (v/v) FBS) at a final concentration of 5 % (v/v) DMSO. Place cryopreservation medium on ice until ready to use.

3. Aseptically remove medium from the flask(s).

4. Add appropriate volumes of sterile Ca²⁺- and Mg²⁺-free DPBS to the side of the flask so as to avoid dislodging the cells (see Table 3).

5. Rinse the cells with DPBS (using a gently rocking motion) and discard.

6. Add appropriate volume of 0.25 % (w/v) Trypsin-0.53 mM EDTA solution to the flask (see Table 3).

7. Incubate the flask at 37 °C ± 1 °C until the cells round up. Observe cells under an inverted microscope every 5 min. When the flask is tilted, the attached cells should slide down the surface. This usually occurs after 5 min to 10 min of incubation.

   *Note: Do not leave trypsin-EDTA on the cells any longer than necessary as clumping may result.*

8. Neutralize the trypsin-EDTA by adding an equal volume of complete growth medium (EMEM + 10 % (v/v) FBS) to each flask. Disperse the cells by pipetting, gently, over the surface of the monolayer. Pipette the cell suspension up and down with the tip of the pipette resting on the bottom corner or edge until a single cell suspension is obtained. Care should be taken to avoid the creation of foam.

9. Using a 1 mL pipette, remove 1 mL of cell suspension for total cell count and viability.

10. Record total cell count and viability.

11. Spin cells at approximately 125 xg for 5 min to 10 min at room temperature. Carefully aspirate and discard the medium, leaving the pellet undisturbed.

12. Calculate volume of cryopreservation medium based on the count performed at step 9 and resuspend pellet in cold cryopreservation medium at a viable cell density of 2 x 10⁶ (acceptable range: 1.0 x 10⁶ cells/mL to 2.0 x 10⁶ cells/mL). by gentle pipetting up and down.

13. Dispense 1 mL of cell suspension, using a 5 mL or 10 mL pipette, into each 1.0 mL cryovial.

14. Place filled cryovials at 2 °C to 8 °C until ready to cryopreserve. A minimum equilibration time of 10 min but no longer than 45 min is necessary to allow DMSO to penetrate the cells.

   *Note: DMSO is toxic to the cells. Long exposure in DMSO may affect viability.*
6. Cryopreservation of Cells

Material:

- Liquid nitrogen freezer
- Cryomed Programmable freezer (Forma Scientific, catalog no. 1010) or
- Mr. Frosty (Nalgene, catalog no. 5100)
- Isopropanol
- Cryovial rack

a. Cryopreservation using a rate-controlled programmable freezer

Method:

A slow and reproducible cooling rate is very important to ensure good recovery of cultures. A decrease of −1 °C per min to -80 °C followed by rapid freeze at about 15 °C to 30 °C per min drop to -150 °C will usually work for most animal cell cultures. The best way to control the cooling process is to use a programmable, rate-controlled, electronic freezer unit. Refer to the manufacturer’s handbook for detailed procedure.

i. Using the Cryomed

Starting the Cryopreservation Process

1. Check that the liquid nitrogen valve that supplies the Cryomed is open.

2. Check the gauge to ensure that there is enough liquid nitrogen in the open tank to complete the freeze.

3. Install the thermocouple probe so that the tip is immersed midway into the control fluid

   **Note:** Be sure that the thermocouple is centered in the vial and the vial is placed centered in the rack. The probe should be changed after three uses or if it turns yellow to ensure accurate readings by the controller during the freezing process. Old medium may have different freezing characteristics.

4. Close and latch Cryomed door.

5. Turn on microcomputer, computer and monitor.

6. Double click the “Cryomed” icon. The machine may need to be pre-programmed for specific cell type and medium.

7. From the top of the screen, select MENU → RUN FUNCTIONS → START RUN.
8. Fill out the box which appears on the screen. Cell line ID; TYPE OF SAMPLE; MEDIA; NUMBER OF SAMPLES.

9. Hit the ESCAPE key and the Cryomed will cool to 4°C.

10. Once Cryomed chamber has cooled to 4°C, load cryovials onto racks and close the door.

11. When the Cryomed’s chamber temperature and the sample temperature have reached approximately 4°C; press the space bar to initiate the rate controlled cryopreservation process.

Completing the Cryopreservation Process

1. When samples have reached –80°C, an alarm will sound. To silence this, select ALARM from the options at the top of the screen.

2. Select MENU → RUN FUNCTIONS → STOP. Hit the ESCAPE key to return to the main menu and select EXIT.

3. Immediately transfer vials to liquid nitrogen freezer.

4. Shut down the microcomputer and then turn off the monitor.

b. Cryopreservation using “Mr. Frosty”

1. One day before freezing cells, add 250 mL isopropanol to the bottom of the container and place at 2 °C to 8 °C.

2. On the day of the freeze, prepare cells for cryopreservation as described above.

3. Insert cryovials with the cell suspension in appropriate slots in the container.

4. Transfer the chamber to a –70 °C to –90 °C freezer and store overnight.

5. Next day, transfer cryovials to the vapor phase of liquid nitrogen freezer.

Note: Each container has 18 slots which can accommodate 18 cryovials in one freeze.
Important information when using the rate-controlled programmable freezer or a manual method (Mr. Frosty) for cryopreservation of mammalian cells.

- Regardless which cooling method is used, it is important that the transfer to the final storage location (between -130 °C and –196 °C) be done quickly and efficiently. If the transfer cannot be done immediately, the vials can be placed on dry ice for a short time. This will avoid damage to cultures by inadvertent temporary warming during the transfer process. Warming during this transfer process is a major cause of variation in culture viability upon thawing.

- Always keep the storage temperature below –130 °C for optimum survival. Cells may survive storage at higher temperatures but viability will usually decrease over time. The ideal storage container is a liquid nitrogen freezer where the cultures are stored in the vapor phase above the liquid nitrogen.

*Note: ATCC does not have experience in the cryopreservation of the U-87 MG cells by any other method than the Cryomed programmable freezer.*

7. Storage

- Store cryopreserved cells in the vapor phase of liquid nitrogen freezer (below –130 °C) for optimum long-term survival.

*Note: Experiments on long-term storage of animal cell lines at different temperature levels indicate that a -70 °C storage temperature is not adequate except for very short period of time. A -90 °C storage may be adequate for longer periods depending upon the cell line preserved. The efficiency of recovery, however, is not as great as when the cells are stored in vapor phase of the liquid nitrogen freezer.*
APPENDIX 1: PHOTOMICROGRAPHS OF NCI-PBCF-HTB14 (U-87 MG)

Figure 1: Photomicrograph of U-87 MG cells after one day, post-freeze recovery. Cells were plated at 5 x 10^4 viable cells/cm^2.

Figure 2: Photomicrograph of U-87 MG cells after two days, post-freeze recovery. Cells were plated at 5 x 10^4 viable cells/cm^2.
Figure 3: Photomicrograph of U-87 MG cells at various time points after seeding at a cell density of $4 \times 10^4$ viable cells/cm$^2$. 
**APPENDIX 2: GROWTH PROFILE FOR NCI-PBCF-HTB14 (U-87 MG)**

![Growth profile graph](image)

**Figure 3:** Growth profile of U-87 MG cells; cells were plated at $4 \times 10^4$ viable cells/cm²; population doubling time (PDT) is approximately 34 h.
# APPENDIX 3: CYTOGENETIC ANALYSIS OF NCI-PBCF-HTB14 (U-87 MG) CELLS

## NCI-PBCF-HTB14 Karyotype Results

<table>
<thead>
<tr>
<th>Metaphase Spread</th>
<th>Karyotype</th>
</tr>
</thead>
<tbody>
<tr>
<td><img src="image" alt="Metaphase Spread Image" /></td>
<td><img src="image" alt="Karyotype Image" /></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Number of metaphase spreads counted</th>
<th>20</th>
</tr>
</thead>
<tbody>
<tr>
<td>Band level</td>
<td>300-400</td>
</tr>
<tr>
<td>Number of metaphase spreads karyotyped</td>
<td>10</td>
</tr>
<tr>
<td>Chromosome range</td>
<td>43-45</td>
</tr>
<tr>
<td>Sex</td>
<td>Male</td>
</tr>
<tr>
<td>Comments</td>
<td>Aneuploid*</td>
</tr>
</tbody>
</table>

**Karyotype:**

43-45,X,-1,add(1)(p13)x2,add(6)(p21.1),add(6)(q21),+9,del(9)(p13),der(10;16)(q10;p10), -11,-12,-13,-14,-16,del(18)(q21.3),-20,+mar1,+mar2,+mar3,+mar4,+mar5,+1-3mar  
(*ISCN nomenclature written based on a diploid karyotype*).

*Human diploid karyotype (2N): 46,XX (female) or 46,XY (male)*
Karyotype Summary:
In the karyotype image, arrows indicate regions of abnormality. It should be noted that the karyotype description includes the observed abnormalities from all examined metaphase spreads, but due to heterogeneity, not all of the karyotyped cells will contain every abnormality.

This is a highly rearranged human cell line of male origin containing 43 to 45 chromosomes per metaphase spread (hypodiploid). Structural abnormalities include rearrangements to chromosomes 1, 6, 9, 10, 16 and 18. There are five unidentifiable clonal marker chromosomes (markers present in two or more of the examined cells) [+mar1 through +mar5] and one to three unidentifiable non-clonal marker chromosomes (markers present in only one of the examined cells).

The rearrangements include:
- Addition of unknown material to the short arms (designated by p) of chromosomes 1 and 6;
- Addition of unknown material to the long arm (designated by q) of chromosome 6;
- Deletion of material from the short arm of chromosome 9;
- Deletion of material from the long arm of chromosome 18;
- A translocation involving chromosomes 10 and 16 [der(10;16)(q10;p10)];

Numerical changes are based on a diploid karyotype which would contain two copies of each chromosome (2N). Therefore, karyotype designations such as -1 and -11, indicates one copy of structurally normal chromosomes 1 and 11; +9 indicates three copies of chromosome 9. (ISCN 2009: An International System for Human Cytogenetic Nomenclature (2009), Editors: Lisa G. Shaffer, Marilyn L. Slovak, Lynda J. Campbell)
Karyotype Procedure:

- **Cell Harvest:** Cells were allowed to grow to 80-90% confluence. Mitotic division was arrested by treating the cells with KaryoMax® colcemid for 20 minutes to 2 hours at 37°C. Cells were harvested using 0.05% Trypsin-EDTA, treated with 0.075M KCL hypotonic solution, and then fixed in three changes of a 3:1 ratio of methanol:glacial acetic acid.

- **Slide Preparation:** Slides were prepared by dropping the cell suspension onto wet glass slides and allowing them to dry under controlled conditions.

- **G-banding:** Slides were baked one hour at 90°C, trypsinized using 10X trypsin-EDTA, and then stained with Leishman's stain.

- **Microscopy:** Slides were scanned using a 10X objective and metaphase spreads were analyzed using a 100X plan apochromat objective on an Olympus BX-41 microscope. Imaging and karyotyping were performed using Cytovision® software.

- **Analysis:** Twenty metaphase cells were counted and analyzed, and representative metaphase cells were karyotyped depending on the complexity of the study.

Summary of Karyotyping Procedure:

G-band karyotyping analysis is performed using GTL banding technique: G bands produced with trypsin and Leishman. Slides prepared with metaphase spreads are treated with trypsin and stained with Leishman's. This method produces a series of light and dark bands that allow for the positive identification of each chromosome.

**U-87 MG cell line** karyotyping was carried out by Cell Line Genetics, Inc. (Madison, WI 53719)
APPENDIX 4: GLOSSARY OF TERMS

**Confluent monolayer:** adherent cell culture in which all cells are in contact with other cells all around their periphery and no available substrate is left uncovered.

**Split ratio:** the divisor of the dilution ration of a cell culture to subculture (e.g., one flask divided into four, or 100 mL up to 400 mL, would be split ratio of 1:4).

**Subculture (or passage):** the transfer or transplantation of cells, with or without dilution, from one culture vessel to another.

**Passage No:** the total number of times the cells in the culture have been subcultured or passaged (with each subculture the passage number increases by 1).

**Population doubling level (PDL):** the total number of population doublings of a cell line since its initiation in vitro (with each subculture the population doubling increases in relationship to the split ratio at which the cells are plated). See Appendix 7.

**Population doubling time (doubling time):** the time interval, calculated during the logarithmic phase of growth in which cells double in number.

**Seeding density:** recommended number of cells per cm² of substrate when inoculating a new flask.

**Epithelial-like:** adherent cells of a polygonal shape with clear, sharp boundaries between them.

**Fibroblast-like:** adherent cells of a spindle or stellate shape.
APPENDIX 5: REFERENCE


APPENDIX 6: REAGENT LOT TRACEABILITY AND CELL EXPANSION TABLES

Table 4: Reagent Lot Traceability

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<th>Lot #</th>
<th>Expiration Date</th>
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Table 5: Cell Expansion

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<th>CELL COUNT</th>
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APPENDIX 7: CALCULATION OF POPULATION DOUBLING LEVEL (PDL)

Calculate the PDL of the current passage using the following equation:

$$\text{PDL} = X + 3.322 (\log Y - \log I)$$

Where:
- $X$ = initial PDL
- $I$ = cell inoculum (number of cells plated in the flask)
- $Y$ = final cell yield (number of cells at the end of the growth period)

APPENDIX 8: SAFETY PRECAUTIONS

- Use at least approved Biological Safety Level 2 (BSL-2) facilities and procedures.
- Wear appropriate Personal Protective Equipment (PPE), such as isolation gown, lab coat with sleeve protectors, face shield and gloves.
- Use safety precautions for working with liquid nitrogen, nitrogen vapor, and cryogenically cooled fixtures apply.
- Use liquid nitrogen freezers and liquid nitrogen tanks only in areas with adequate ventilation. Liquid nitrogen reduces the concentration of oxygen and can cause suffocation.
- Wear latex gloves over insulating gloves to prevent liquid nitrogen from soaking in and being held next to the skin. Liquid nitrogen is extremely cold and will cause burns and frostbite. Metal inventory racks, tank components, and liquid nitrogen transfer hoses exposed to liquid nitrogen or nitrogen vapor quickly cool to cryogenic temperatures and can cause burns and frostbite.
- Wear a full face mask when thawing and retrieving vials from liquid nitrogen freezer. Danger to the technician derives mainly from the possibility that liquid nitrogen can penetrate the cryovial during storage. On warming, rapid evaporation of the nitrogen within the confines of such cryovial can cause an aerosol or explosion of the cryovial and contents.