

144	T75	75 cm ² tissue culture flask
145	T150	150 cm ² tissue culture flask
146		

146

LIST OF FIGURES

147	Figure 7-1	pGudLuc7.ERE Plasmid.....	8
148	Figure 9-1	Hemocytometer Counting Grid.....	20
149	Figure 11-1	Example Concentration Curve for Calculation of Relative EC ₅₀ Values.....	35
150	Figure 11-2	Example Concentration Curve for Calculation of Absolute EC ₅₀ Values.....	36
151	Figure 12-1	Agonist Range Finder Test Plate Layout.....	37
152	Figure 13-1	Agonist Comprehensive Test Plate Layout.....	39
153	Figure 14-1	Example Scatter Chart of the Methoxychlor Control QC.....	41
154	Figure 14-2	Example Figure of a Sigmoidal E2 Concentration Response Curve.....	42
155	Figure 14-3	Example Scatter Chart of the E2 EC ₅₀ Control QC.....	43
156	Figure 14-4	Example Scatter Chart of the DMSO Control QC.....	44
157			

157

LIST OF TABLES

158	Table 6-1	Concentration of E2 Reference Standard Used in Range Finder and	
159		Comprehensive Testing.....	6
160	Table 10-1	Preparation of E2 Stock Solution	24
161	Table 10-2	Preparation of E2 Reference Standard Dosing Solution for Range	
162		Finder Testing.....	25
163	Table 10-3	Preparation of Test Substance Dosing Solutions for Range Finder	
164		Testing.....	25
165	Table 10-4	Preparation of Test Substance Dosing Solutions for Comprehensive	
166		Testing.....	26
167	Table 11-1	Visual Observation Scoring.....	30
168			

169 **1.0 PURPOSE**

170 This protocol is designed to evaluate coded test substances for potential estrogen receptor (ER)
171 agonist activity using the LUMI-CELL® ER assay.

172 **2.0 SPONSOR**

173 The National Toxicology Program (NTP) Interagency Center for the Evaluation of Alternative
174 Toxicological Methods (NICEATM), P.O. Box 12233 Research Triangle Park, NC 27709

175

176 William S. Stokes, DVM, DACLAM

177 Rear Admiral, U.S. Public Health Service

178 Chief Veterinary Officer, USPHS

179 Director, NICEATM

180 National Institute of Environmental Health Sciences, NIH, DHHS

181 Bldg. 4401, Room 3129, MD EC-14

182 79 T.W. Alexander Drive

183 Research Triangle Park, NC 27709

184 Phone: 919-541-7997

185 Fax: 919-541-0947

186 Email: stokes@niehs.nih.gov

187

188 Raymond Tice, Ph.D.

189 Deputy Director, NICEATM

190 National Institute of Environmental Health Sciences

191 MD EC-17, P.O. Box 12233

192 Research Triangle Park, NC 27709

193 Phone: 919-541-4482
194 FAX: 919-541-0947
195 Email: tice@niehs.nih.gov
196
197 David Allen, Ph.D.
198 Principal Investigator
199 ILS, Inc./Contractor supporting NICEATM
200 National Institute of Environmental Health Sciences
201 MD EC-17, P.O. Box 12233
202 Research Triangle Park, NC 27709
203 Phone: 919-316-4587
204 FAX: 919-541-0947
205 Email: allen7@niehs.nih.gov
206
207 Frank Deal, M.S.
208 Staff Toxicologist
209 ILS, Inc./Contractor supporting NICEATM
210 National Institute of Environmental Health Sciences
211 MD EC-17, P.O. Box 12233
212 Research Triangle Park, NC 27709
213 Phone: 919-316-4587
214 FAX: 919-541-0947
215 Email: dealf@niehs.nih.gov
216
217

217 Patricia Ceger, M.S.
218 Project Coordinator/Technical Writer
219 ILS, Inc./Contractor supporting NICEATM
220 National Institute of Environmental Health Sciences
221 MD EC-17, P.O. Box 12233
222 Research Triangle Park, NC 27709
223 Phone: 919-316-4556
224 Fax: 919-541-0947
225 E-Mail: cegerp@niehs.nih.gov

226

227 **2.1 Substance Inventory and Distribution Management**

228 Cynthia Smith, Ph.D.
229 Chemistry Resources Group Leader
230 National Institute of Environmental Health Sciences
231 MD EC-06, P.O. Box 12233
232 Research Triangle Park, NC 27709
233 Phone: 919-541-3473

234

235 **3.0 DEFINITIONS**

- 236 • **Dosing Solution:** The test substance, control substance, or reference standard
237 solution, which is to be placed into the tissue culture wells for experimentation.
- 238 • **Raw Data:** Raw data includes information that has been collected but not
239 formatted or analyzed, and consists of the following:
- 240 ○ Data recorded in the Study Notebook

- 241 ○ Computer printout of initial luminometer data
- 242 ○ Other data collected as part of GLP compliance, e.g.:
- 243 ▪ Equipment logs and calibration records
- 244 ▪ Test substance and tissue culture media preparation logs
- 245 ▪ Cryogenic freezer inventory logs
- 246 • **Soluble:** Test substance exists in a clear solution without visible cloudiness or
- 247 precipitate.
- 248 • **Study Notebook:** The study notebook contains recordings of all activities related
- 249 to the conduct of the LUMI-CELL® ER TA agonist assay.
- 250 • **Test Substances:** Substances supplied to the testing laboratories that are coded
- 251 and distributed such that only the Project Officer, Study Management Team
- 252 (SMT), and the Substance Inventory and Distribution Management have
- 253 knowledge of their true identity. The test substances will be purchased, aliquoted,
- 254 coded, and distributed by the Supplier under the guidance of the NIEHS/NTP
- 255 Project Officer and the SMT.

256

257 4.0 **TESTING FACILITY AND KEY PERSONNEL**

258 4.1 **Testing Facility**

259 Xenobiotic Detection Systems, Inc. (XDS), 1601 E. Geer St., Suite S, Durham, NC 27704

260 4.2 **Key Personnel**

- 261 • Study Director: John Gordon, Ph.D.
- 262 • Laboratory Technician(s): Cynthia Matherly
- 263 • Scientific Advisor: Mike Denison, Ph.D.
- 264 • Quality Assurance Director: Andrew Chu
- 265 • Safety Manager: Andrew Chu

- 266 • Facility Management: George Clark, Ph.D., Robert Clark, MS, Tina Ginter,
267 Andrew Chu
- 268 • Laboratory Director: Andrew Chu

269 5.0 IDENTIFICATION OF TEST AND CONTROL SUBSTANCES

270 5.1 Test Substances

271 Test substances are coded and will be provided to participating laboratories by the Substance
272 Inventory and Distribution Management team.

273 5.2 Controls

274 Controls for the ER agonist protocol are as follows:

275 *Vehicle control (dimethyl sulfoxide [DMSO]):* 1% (v/v) DMSO (CASRN 67-68-5) diluted in
276 tissue culture media.

277 *Reference standard (17β-estradiol [E2]):* Three concentrations of E2 (CASRN 50-28-2) in
278 duplicate for range finder testing and a serial dilution consisting of 10 concentrations of E2 in
279 duplicate for comprehensive testing

280 *Positive control (p,p'-Methoxychlor [methoxychlor]):* Methoxychlor (CASRN 72-43-5), 3.13
281 µg/mL in tissue culture media, used as a weak positive control.

282 6.0 OVERVIEW OF GENERAL PROCEDURES FOR AGONIST TESTING

283 All experimental procedures are to be carried out under aseptic conditions and all solutions,
284 glassware, plastic ware, pipettes, etc., shall be sterile. All methods and procedures shall be
285 documented in the study notebook.

286 Agonist range finder testing is conducted on 96-well plates using three concentrations (1.00×10^{-4} ,
287 2.00×10^{-6} , 1.00×10^{-7} µg/mL) of E2 in duplicate as the reference standard and six replicate
288 wells for the DMSO control. Range finder testing uses all wells of the 96-well plate.

289 Comprehensive testing is conducted on 96-well plates using 10 concentrations of E2 in duplicate
290 as the reference standard (**Table 6-1**). Four replicate wells for the DMSO control and three

291 replicate wells for the methoxychlor control are included on each plate. To avoid edging effects¹,
292 wells on the perimeter of the plate are not used for experiments. These wells should contain
293 tissue culture media only (i.e., no cells).

294 **Table 6-1 Concentrations of E2 Reference Standard Used in Comprehensive Testing**

E2 Concentrations ¹		
1.00 x 10 ⁻⁴	6.25 x 10 ⁻⁶	1.95 x 10 ⁻⁷
5.00 x 10 ⁻⁵	3.13 x 10 ⁻⁶	9.78 x 10 ⁻⁸
2.50 x 10 ⁻⁵	1.56 x 10 ⁻⁶	
1.25 x 10 ⁻⁵	7.83 x 10 ⁻⁷	

295 ¹Concentrations are presented in µg/mL.

296 Visual observations for cell viability are conducted for all experimental plates just prior to
297 LUMI-CELL® ER evaluation. CellTiter-Glo® based cell viability testing (when used) is
298 conducted concurrently in parallel plates, as outlined in **Section 11.2**.

299 Luminescence data, measured in relative light units (RLUs), is corrected for background
300 luminescence by subtracting the mean RLU value of the vehicle control (DMSO) wells from the
301 RLU measurements for each of the other wells of the 96-well plate. Data is then transferred into
302 Excel® data management spreadsheets and GraphPad PRISM® 4.0 statistical software, graphed,
303 and evaluated as follows:

- 304 • A response is considered positive for agonist activity when the average adjusted
305 RLU for a given concentration is greater than the mean RLU value plus three
306 times the standard deviation for the vehicle control.
- 307 • Any response below this threshold is considered negative for agonist activity.

308 For substances that are positive at one or more concentrations, the concentration that causes a
309 half-maximal response (EC₅₀) is calculated using a Hill function analysis. The Hill function is a
310 four-parameter logistic mathematical model relating the substance concentration to the response
311 (typically following a sigmoidal curve) using the equation below:

¹ Edging effects are variations in response seen in the outermost wells in a cell culture plate. These variations are believed to be due to variations in temperature, evaporation, etc., that may occur in these wells that would ultimately affect cellular growth and health (ICCVAM, 2001).

312
$$Y = \text{Bottom} + \frac{\text{Top} - \text{Bottom}}{1 + 10^{(\log EC_{50} - X) \text{HillSlope}}}$$

313 where Y = response (i.e., relative light units); X = the logarithm of concentration; Bottom = the
314 minimum response; Top = the maximum response; log EC₅₀ = the logarithm of X as the response
315 midway between Top and Bottom; and HillSlope describes the steepness of the curve. The model
316 calculates the best fit for the Top, Bottom, HillSlope, and EC₅₀ parameters. See **Section 11.6.5**
317 for more details.

318 Acceptance or rejection of a test is based on evaluation of reference standard and control results
319 from each experiment conducted on a 96-well plate. Results for these controls are compared to
320 historical results compiled in the historical database, as seen in **Section 14.0**.

321 **6.1 Range Finder Testing**

322 Agonist range finding for coded substances consists of a six point, logarithmic serial dilution
323 using duplicate wells per concentration. Concentrations for comprehensive testing are selected
324 based on the response observed in range finder testing. If necessary, a second range finder test
325 can be conducted to clarify the optimal concentration range to test (see **Section 12.0**).

326 **6.2 Comprehensive Testing**

327 Comprehensive agonist testing for coded substances consists of 11 point, double serial dilutions,
328 with each concentration tested in triplicate wells of the 96-well plate. Three separate experiments
329 are conducted for comprehensive testing on three separate days, except during Phases III and IV
330 of the validation effort, in which comprehensive testing experiments are conducted once (see
331 **Section 13.0**).

332 **7.0 MATERIALS FOR LUMI-CELL® ER AGONIST TESTING**

333 This section provides the materials needed to conduct LUMI-CELL® testing, with associated
334 brand names/vendors² in brackets.

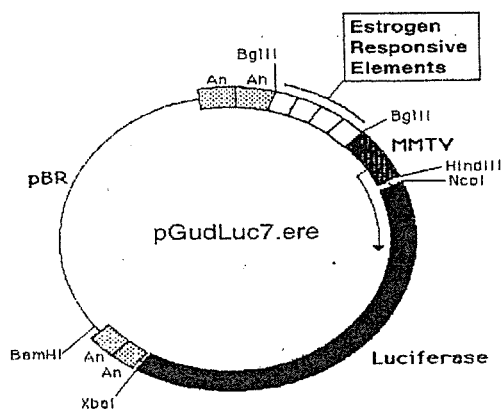
335

²Brand names and vendors should not be considered an endorsement by the U.S. Government or any member of the U.S. Government; such information is provided as examples.

335 **7.1 BG1Luc4E2 Cells:**

336 Human ovarian cancer cell line stably transfected with a plasmid containing an estrogen response
337 element pGudLuc7.0 (Figure 7-1) [XDS].

338 **Figure 7-1 pGudLuc7.ERE Plasmid.**



339

340 **7.2 Technical Equipment:**

341 All technical equipment may be obtained from Fisher Scientific International, Inc. (Liberty Lane
342 Hampton, NH, USA 03842). Equivalent technical equipment from another commercial source
343 can be used.

- 344 • Analytical balance (Cat. No. 01-910-320)
- 345 • Berthold Orion 1 Microplate Luminometer [Berthold CatNo.: Orion 1 MPL3] or
346 equivalent and dedicated computer
- 347 • Biological safety hood, class II, and stand (Cat. No. 16-108-99)
- 348 • Centrifuge (low speed, tabletop with swinging bucket rotor) (Cat. No. 04-978-50
349 centrifuge, and 05-103B rotor)
- 350 • Combustion test kit (CO₂ monitoring) (Cat. No. 10-884-1)
- 351 • Drummond diaphragm pipetter (Cat. No. 13-681-15)
- 352 • Freezers, -20°C (Cat. No. 13-986-150), and -70°C (Cat. No. 13-990-86)
- 353 • Hand tally counter (Cat. No. 07905-6)

- 354 • Hemocytometer, cell counter (Cat. No. 02-671-5)
- 355 • Light microscope, inverted (Cat. No. 12-561-INV)
- 356 • Light microscope, upright (Cat. No. 12-561-3M)
- 357 • Liquid nitrogen flask (Cat. No. 11-675-92)
- 358 • Micropipetter, repeating (Cat. No. 21-380-9)
- 359 • Pipetters, air displacement, single channel (0.5 –10µl (Cat. No. 21-377-191), 2 –
- 360 20 µl (Cat. No. 21-377-287), 20 – 200 µl (Cat. No. 21-377-298), 200 - 1000 µl
- 361 (Cat. No. 21-377-195))
- 362 • Refrigerator/freezer (Cat. No. 13-986-106A)
- 363 • Shaker for 96-well plates (Cat. No. 14-271-9)
- 364 • Sodium hydroxide (Cat. No. 5318-500)
- 365 • Sonicating water bath (Cat. No. 15-335-30)
- 366 • Tissue culture incubator with CO₂ and temperature control (Cat. No. 11-689-4)
- 367 • Vacuum pump with liquid trap (side arm Erlenmeyer) (Cat. No. 01-092-29)
- 368 • Vortex mixer (Cat. No. 12-814)

369 Equipment should be maintained and calibrated as per GLP guidelines and individual laboratory
370 SOPs.

371 **7.3 Reference Standard, Controls, and Tissue Culture Supplies**

372 All tissue culture reagents must be labeled to indicate source, identity, storage conditions and
373 expiration dates. Tissue culture solutions must be labeled to indicate concentration, stability
374 (where known), and preparation and expiration dates.

375 Equivalent tissue culture media and sera from another commercial source can be used, but must
376 first be tested as described in **Section 15.0** to determine suitability for use in this test method.

377 The following are the necessary tissue culture reagents and possible commercial sources (in
378 brackets) based on their use in the pre-validation studies:

- 379 • 17 β -estradiol (CAS RN: 50-28-2) [Sigma-Aldrich, Cat. No. E8875]
- 380 • CellTiter-Glo® Luminescent Cell Viability Assay [Promega Cat. No. G7572]
- 381 • Cryovial, 2 mL (Corning Costar) [Fisher Scientific Cat. No. 03-374-21]
- 382 • Culture tube 13 x 100mm (case) [Thomas Scientific Cat. No.: 10009186R38]
- 383 • Culture tube, 15 mL conical (Corning Costar) [Fisher Scientific Cat. No. 05-538-
- 384 51]
- 385 • Culture tube, 50 mL conical (Corning Costar) [Fisher Scientific Cat. No. 05-
- 386 526C]
- 387 • DMSO, U.S.P. analytical grade. [Sigma-Aldrich, Cat. No. 34869-100ML]
- 388 • Dulbecco's Modification of Eagle's Medium (DMEM), containing 4.5 g/L
- 389 glucose, with sodium pyruvate, without phenol red or L-glutamine
- 390 [Mediatech/Cellgro, Cat. No. 17-205-CV]
- 391 • Fetal Bovine Serum [Mediatech/Cellgro Cat. No. MT 35-010-CV]
- 392 • Fetal Bovine Serum, charcoal/dextran treated, triple 0.1 μ m sterile filtered
- 393 [Hyclone, Cat. No. SH30068.03]
- 394 • Gentamycin Sulfate (G418), 50 mg/mL [Mediatech/Cellgro Cat. No. 30-234-CR]
- 395 • L-glutamine, 29.2 mg/mL [Cellgro, Cat. No. 25005-CI]
- 396 • Luciferase Assay System (10-Pack) [Promega Cat. No. E1501]
- 397 • Lysis Solution 5X [Promega, Cat. No. E1531]
- 398 • Methoxychlor (CAS RN: 72-43-5) [Sigma-Aldrich, Cat. No. 49054]
- 399 • Penicillin/streptomycin solution, 5000 I.U. penicillin, 5000 μ g/mL streptomycin
- 400 [Cellgro, Cat. No. 30-001-CI].
- 401 • Phosphate buffered saline (PBS, 1X) without calcium and magnesium [Cellgro,
- 402 Cat. No. 21-040-CV]
- 403 • Pipettes, serological: 2.0 mL [Sigma-Aldrich, Cat. No. P1736], 5.0 mL [Sigma-
- 404 Aldrich, Cat. No. P1986], 25 mL [Sigma-Aldrich, Cat. No. P2486]

- 405 • RPMI 1640 medium, containing L-glutamine [Mediatech, Cat. No. 10-040-CV]
- 406 • Tissue culture flasks (Corning-Costar): 25 cm² (T25) [Fisher Cat. No. 10-126-28];
- 407 75 cm² (T75) [Fisher Cat. No. 10-126-37]; and 150 cm² (T150) [Fisher Cat. No.
- 408 10-126-34]
- 409 • Tissue culture plates (Corning-Costar): 96-well [Thomas Scientific Cat. No.
- 410 6916A05]
- 411 • Trypsin (10X), 2.5% in Hank's balanced salt solution (HBSS), without calcium
- 412 and magnesium, without phenol red [Cellgro, Cat. No. 25-054-CI].

413 All reagent lot numbers and expiration dates must be recorded in the study notebook.

414 **8.0 PREPARATION OF TISSUE CULTURE MEDIA AND SOLUTIONS**

415 All tissue culture media and media supplements must be quality tested before use in experiment
416 (see Section 15.0).

417 **8.1 RPMI 1640 Growth Medium (RPMI)**

418 RPMI 1640 is supplemented with 0.9% Pen-Strep and 8.0% FBS to make RPMI growth medium
419 (RPMI).

420 Procedure for one 549 mL bottle:

- 421 1. Remove FBS from -70°C freezer, and Pen-Strep from -20°C freezer and allow to
- 422 equilibrate to room temperature.
- 423 2. Add 44 mL of FBS and 5 mL Pen-Strep to the bottle of RPMI 1640.
- 424 3. Label RPMI bottle as indicated in **Section 7.3**

425 *Store at 2-8°C for no longer than six months or until the shortest expiration date of any media*
426 *component.*

427 **8.2 Estrogen-Free DMEM Medium**

428 DMEM is supplemented to contain 4.5% charcoal/dextran treated FBS, 1.9% L-glutamine, 0.9%
429 Pen-Strep.

430 Procedure for one 539 mL bottle:

- 431 1. Remove charcoal/dextran treated FBS from -70°C freezer, and L-glutamine and
432 Pen-Strep from -20°C freezer and allow to equilibrate to room temperature.
- 433 2. Add 24 mL of charcoal/dextran treated FBS, 10 mL L-glutamine, and 5 mL Pen-
434 Strep to one 500 mL bottle of DMEM.
- 435 3. Label estrogen-free DMEM bottle as indicated in **Section 7.3**
- 436 *Store at 2-8°C for no longer than six months or until the shortest expiration date of any media*
437 *component..*

438 **8.3 1X Trypsin Solution**

439 1X Trypsin solution is prepared by dilution from a 10X premixed stock solution. The 10X stock
440 solution should be stored in 10 mL aliquots in a -20°C freezer.

441 Procedure for making 100 mL of 1X trypsin:

- 442 1. Remove a 10 mL aliquot of 10X trypsin from -20°C freezer and allow to
443 equilibrate to room temperature.
- 444 2. Aliquot 1 mL Trypsin (10X) along with 9 mL of 1X PBS into ten 15 mL sterile
445 centrifuge tubes.
- 446 3. Label 1X trypsin aliquots as indicated in **Section 7.3**

447 *1X Trypsin should be stored at -20°C.*

448 **8.4 1X Lysis Solution**

449 Lysis solution is prepared by dilution from a 5X premixed stock solution. Both the 5X and 1X
450 solutions can be repeatedly freeze-thawed.

451 The procedure for making 10 mL of 1X lysis solution:

- 452 1. Thaw the 5X Promega Lysis solution and allow it to reach room temperature.
- 453 2. Remove 2 mL of 5X solution and place it in a 15 mL conical centrifuge tube.
- 454 3. Add 8 mL of distilled, de-ionized water to the conical tube.
- 455 4. Cap and shake gently until solutions are mixed.

456 Store at -20°C for no longer than 1 year from receipt.

457 **8.5 Reconstituted Luciferase Reagent**

458 Luciferase reagent consists of two components, luciferase buffer and lyophilized luciferase
459 substrate.

460 For long term storage, unopened containers of the luciferase buffer and lyophilized luciferase
461 substrate can be stored at -70°C for up to one year.

462 To reconstitute luciferase reagent:

- 463 1. Remove luciferase buffer and luciferase substrate from -70°C freezer and allow
464 them to equilibrate to room temperature.
- 465 2. Add 10 mL of luciferase buffer solution to luciferase substrate container and swirl
466 or vortex gently to mix; the Luciferase substrate should readily go into solution.
- 467 3. After solutions are mixed, aliquot to a 15mL centrifuge tube.
- 468 4. Store complete solution at -20°C.

469 Reconstituted luciferase reagent is stable for up to 1 month at -20°C.

470 **8.6 Reconstituted CellTiter-Glo® Reagent**

471 CellTiter-Glo® reagent consists of two components, CellTiter-Glo® buffer and lyophilized
472 CellTiter-Glo® substrate.

473 For long term storage, unopened containers of the CellTiter-Glo® buffer and lyophilized
474 CellTiter-Glo® substrate can be stored at -70°C for up to one year.

475 To reconstitute CellTiter-Glo® reagent:

- 476 1. Remove CellTiter-Glo® buffer and CellTiter-Glo® substrate from -70°C freezer
477 and allow them to equilibrate to room temperature.
- 478 2. Add CellTiter-Glo® buffer solution to CellTiter-Glo® substrate container and
479 swirl or vortex gently to mix; the CellTiter-Glo® substrate should readily go into
480 solution.
- 481 3. After solutions are mixed aliquot to a 15mL centrifuge tube.

482 4. Store complete solution at -20°C .

483 Reconstituted CellTiter-Glo® reagent is stable for up to 3 months at -20°C .

484 **9.0 OVERVIEW OF PROPAGATION AND EXPERIMENTAL PLATING OF**
485 **BG1Luc4E2 CELLS**

486 The BG1Luc4E2 (BG-1) cells are stored in liquid nitrogen in 2 mL cryovials. BG-1 cells are
487 grown as a monolayer in tissue culture flasks in a dedicated tissue culture incubator at $37^{\circ}\text{C} \pm$
488 1°C , $90\% \pm 5\%$ humidity, and $5.0\% \pm 1\%$ CO_2/air . The cells should be examined, on a daily
489 basis during working days, under an inverted phase contrast microscope and any changes in
490 morphology and/or adhesive properties must be noted in the study notebook.

491 Two T150 flasks containing cells at 80 to 90% confluence will usually yield a sufficient number
492 of cells to fill four 96-well plates for use in experiments.

493 **9.1 Procedures for Thawing Cells and Establishing Tissue Cultures**

494 Warm all of the tissue culture media and solutions to room temperature by placing them under
495 the tissue culture hood several hours before use.

496 All tissue culture media, media supplements, and tissue culture plasticware must be quality
497 tested before use in experiments (**Section 15.0**).

498 **9.1.1 Thawing Cells**

- 499 1. Remove a cryovial of frozen BG-1 cells from the liquid nitrogen flask.
- 500 2. Facilitate rapid thawing by loosening the top slightly (do not remove top) to
501 release trapped gasses and retightening it. Roll vial between palms.
- 502 3. Use a micropipette to transfer cells to a 50 mL conical centrifuge tube.
- 503 4. Rinse cryovial twice with 1X PBS and add PBS rinse material to the conical tube.
- 504 5. Add 20 mL of RPMI to the conical tube.
- 505 6. Centrifuge at $1000 \times g$ for eight min. If a pellet of cells has not formed, centrifuge
506 for an additional 5 minutes.

- 507 7. Aspirate media from pellet and re-suspend it in 5 mL RPMI, drawing the pellet
508 repeatedly through a 2.0 mL serological pipette to break up any clumps of cells.
- 509 8. Transfer cells to a T25 flask, place them in an incubator (see conditions in
510 **Section 9.0**) and grow to 80% to 90% confluence (approximately 48 to 72 hrs).

511 9.1.2 Establishing Tissue Cultures

512 Once cells have reached 80 to 90% confluence, transfer the cells to a T75 flask by performing,
513 for example, the following steps:

- 514 1. Remove the T25 flask from the incubator.
- 515 2. Aspirate the RPMI, then add 5 mL 1X PBS
- 516 3. Aspirate 1X PBS, then add 1 to 2 mL 1X trypsin to the T25 flask, gently swirling
517 the flask to coat all cells with the trypsin.
- 518 4. Place the flask in an incubator (see conditions in **Section 9.0**) for 5 to 10 min.
- 519 5. Detach cells by hitting the side of the flask sharply against the palm or heel of the
520 hand.
- 521 6. Confirm cell detachment by examination under an inverted microscope. If cells
522 have not detached, return the flask to the incubator for an additional 2 minutes,
523 then hit the flask again.
- 524 7. After cells have detached, add 5 mL PBS, and transfer the suspended cells to a 50
525 mL centrifuge tube. Wash the flask one additional time with 5 mL PBS.
- 526 8. Immediately add 20 mL RPMI to the conical tube to inhibit further cellular
527 digestion by residual trypsin.
- 528 9. Pellet the cells by centrifugation, as described in **Section 9.1.1**, and re-suspend the
529 cells in 10 mL RPMI medium.
- 530 10. Draw the pellet repeatedly through a 25 mL serological pipette to break up
531 clumps of cells
- 532 11. Transfer cells to a T75 flask, then place the flask in an incubator (see conditions
533 in **Section 9.0**) and grow to 80% to 90% confluence (approximately 48 to 72 hrs).

- 534 When cells have reached 80% to 90% confluency, transfer them into a T150 flask by performing,
535 for example, the following steps:
- 536 12. Remove the T75 flask from the incubator, aspirate the old media and add 5 mL
537 1X PBS.
 - 538 13. Aspirate 1X PBS, add 2 mL of 1X trypsin to the flask, and place it in an incubator
539 (see conditions in **Section 9.0**) for 5 to 10 min.
 - 540 14. Repeat steps **5** through **11** in **Section 9.1.2**, re-suspending the pellet in 20 mL of
541 RPMI.
 - 542 15. Transfer cells to a T150 flask and place it in the incubator (see conditions in
543 **Section 9.0**) and grow to 80% to 90% confluence (approximately 48 to 72 hrs).
 - 544 16. Remove the T150 flask from the incubator.
 - 545 17. Aspirate the RPMI and add 5 mL 1X PBS.
 - 546 18. Aspirate 1X PBS and add 3 mL 1X trypsin to the T150 flask, making sure that the
547 cells are coated with the trypsin.
 - 548 19. Incubate cells in an incubator (see conditions in **Section 9.0**) for 5 to 10 min.
 - 549 20. Detach cells by hitting the side of the flask sharply against the palm or heel of the
550 hand.
 - 551 21. Confirm cell detachment by examination under an inverted microscope. If cells
552 have not detached, return the flask to the incubator for an additional 2 minutes,
553 then hit the flask again.
 - 554 22. After cells have detached, add 5 mL 1X PBS and transfer the suspended cells
555 from the T150 flask to a 50 mL conical tube. Add an additional 5 mL PBS to the
556 flask, swirl around the flask, and then transfer the PBS to the 50 mL conical tube.
 - 557 23. Immediately add 20 mL RPMI to the conical tube to inhibit further cellular
558 digestion by residual trypsin.
 - 559 24. Centrifuge at 1000 x g for eight min. If a pellet of cells has not formed, centrifuge
560 for an additional 5 minutes.

561 25. Aspirate the media from the pellet and re-suspend it in 40 mL RPMI, drawing the
562 pellet repeatedly through a 25 mL serological pipette to break up any clumps of
563 cells.

564 26. Transfer 20 mL of cell suspension to each of two T150 flasks, place them in an
565 incubator (see conditions in **Section 9.0**) and grow to 80% to 90% confluence
566 (approximately 48 to 72 hrs).

567 **9.2 Ongoing Tissue Culture Maintenance, Conditioning in Estrogen-free Medium,**
568 **and Plating Cells for Experimentation**

569 The following procedure is used to condition the BG1Luc4E2 cells to an estrogen-free
570 environment prior to plating the cells in 96-well plates for analysis of estrogen dependent
571 induction of luciferase activity.

572 To start the tissue culture maintenance and estrogen-free conditioning, split the two T150 culture
573 flasks into four T150 flasks. Two of these flasks will be used for continuing tissue culture and
574 will use the RPMI media mentioned above. The other two flasks will be cultured in estrogen-free
575 DMEM for experimental use. Extra care must be taken to avoid contaminating the estrogen-free
576 cells with RPMI.

- 577 1. Remove both T150 flasks from the incubator.
- 578 2. Aspirate the medium and rinse the cells with 5 mL 1X PBS.
- 579 3. Aspirate 1X PBS, then add 3 mL 1X trypsin to the flasks, gently swirling the flask
580 to coat all cells with the trypsin.
- 581 4. Incubate cells in the incubator (see conditions in **Section 9.0**) for 5 to 10 min.
- 582 5. Detach cells by hitting the side of the flask sharply against the palm or heel of the
583 hand.
- 584 6. Confirm cell detachment by examination under an inverted microscope. If cells
585 have not detached, return the flask to the incubator for an additional 2 minutes,
586 then hit the flask again.
- 587 7. After cells have detached, add 5 mL 1X PBS to the first T150 flask and transfer
588 the suspended cells to the second T150 flask.