D341 Medulloblastoma Cell Line

1. Source of cells: Duke University Medical Center, surgical resection from a patient with medulloblastoma as described by H Friedman (1988). Requests for D341 cells should be directed to Darrell Bigner (bigne001@mc.duke.edu).

2. Lineage of cells: tumor-derived cancer cells. Cell origin of medulloblastoma is the neuron or a neural precursor cell.

3. Donor information: 3 year old patient, gender unspecified.

4. Karyotype: abnormal, 47, XY, +8, -22, + del(lXp13), i(17q), with double minutes.

5. Media for cell lines: 1X Zinc Option Media (prepared from Gibco Improved MEM Zinc Option Media 5X Concentrate, formula# 86-0194DJ). Each liter of medium is supplemented with 10 mL HEPES and 40 mL 5.5% NaHCO3. Cells are grown in media supplemented with 10% FBS (Gibco Certified FBS, Cat# 16000-044).

6. Culture conditions: Cells grow suspended in a flask and should be incubated at 37° in the presence of 5% CO2. When disaggregated (i.e. during passage) and grown in ZO media D341 cells will eventually form clumps visible to the naked eye (particularly once cells reach a density of ~200K cells/mL). Cells are grown in T-75 or T-225 filter cap flasks set at an angle (i.e., resting on a sheathed 25 mL pipette, in order to increase area of gas exchange) with up to 150 or 400 mL of media, respectively.

7. Cell Line Maintenance: Cells for passage are transferred to 15 or 50 mL conical tubes and centrifuged at 1260 rpm (319 rcf) for 3 min in a standard benchtop centrifuge with swing-out rotor (both from Eppendorf). Media is aspirated and the cells are resuspended at a density of ~50K cells/mL. The cells should reach a density of 200K-250K/mL after 4-5 days and at this point they are ready for passage at a dilution of 1:5. It is not necessary to spin the cells down every time; at higher culture volumes (i.e. >100 mL) the cells are passaged by diluting the cell suspension directly into new media.

8. Cells may be frozen at a density of 10^6 cells/mL in ZO medium supplemented with 50% FBS and 12% DMSO. To freeze cells, pellet cells as if passing, then resuspend in freezing medium and freeze at a rate of 1 °C/min, then store in the vapor phase of liquid nitrogen. Frozen cells are recovered by thawing in a 37 °C water bath, washing once with PBS or ZO medium, and resuspending in 8 mL of 20% ZO medium in a T-25 flask. Once the medium has begun to turn orange, replace with 2 volumes of 10% ZO medium and maintain as above.

9. Cell passage#: unknown

Notes: D341 cells were selected for study due to 1) neural cell origin, 2) generally diploid karyotype as revealed by digital karyotyping, and 3) activation of oncogenic transcriptional networks (Siu et al., 2003; Di et al., 2005).

References:
