

Banco de Células do Rio de Janeiro

Data Sheet

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BCRJ Code:	0294
Cell Line:	ONS-76
Species:	Homo sapiens
Vulgar Name:	Human
Tissue:	Neural/Brain
Cell Type:	Epithelial
Morphology:	Epithelial-Like
Disease:	Medulloblastoma
Growth Properties:	Adherent
Sex:	Female
Age/Ethinicity:	2 Year /
DNA Profile:	AMELOGENINA: X D5S818: 9,10 D13S317: 8,13 D7S820: 11,12 D16S539: 9,10 vWA: 14,16 TH01: 9 TPOX: 8 CSF1PO:10,12
Biosafety:	1
Addtional Info:	POSITIVE FOR 145 Kda AND 200kDA NEUROFILAMENT AND NSE. Negative for GFAP AND S-100 PROTEIN
Culture Medium:	RPMI-1640 medium modified to contain 2 mM L-glutamine, 4500 mg/L glucose and 10% of fetal bovine serum.

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Subculturing:	Remove and discard culture medium. Briefly rinse the cell layer with PBS without calcium and magnesium to remove all traces of serum which contains trypsin inhibitor. Add 2.0 to 3.0 mL of Trypsin-EDTA solution to flask and observe cells under an inverted microscope until cell layer is dispersed (usually within 5 to 15 minutes). Add 6.0 to 8.0 mL of complete growth medium and aspirate cells by gently pipetting. Add appropriate aliquots of the cell suspension into new culture vessels. Incubate cultures at 37°C. NOTE: For more information on enzymatic dissociation and subculturing of cell lines consult Chapter 12 in Culture of Animal Cells, a manual of Basic Technique by R. Ian Freshney, 6th edition, published by Alan R. Liss, N.Y., 2010.
Subculturing Medium Renewal:	2 to 3 times per week
Culture Conditions:	Atmosphere: air, 95%; carbon dioxide (CO2), 5% Temperature: 37°C
Cryopreservation:	95% FBS + 5% DMSO (Dimethyl sulfoxide)

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Thawing Frozen Cells:	 SAFETY PRECAUTION: It is strongly recommended to always wear protective gloves, clothing, and a full-face mask when handling frozen vials. Some vials may leak when submerged in liquid nitrogen, allowing nitrogen to slowly enter the vial. Upon thawing, the conversion of liquid nitrogen back to its gas phase may cause the vial to explode or eject its cap with significant force, creating flying debris. 1. Thaw the vial by gently agitating it in a 37°C water bath. To minimize contamination, keep the O-ring and cap out of the water. Thawing should be rapid (approximately 2 minutes). 2. Remove the vial from the water bath as soon as its contents are thawed and decontaminate it by dipping in or spraying with 70% ethanol. From this point, all operations must be performed under strict aseptic conditions. 3. For cells sensitive to DMSO, it is recommended to remove the cryoprotective agent immediately. Transfer the vial contents to a centrifuge tube containing 9.0 mL of complete culture medium and centrifuge at approximately 125 × g for 5 to 7 minutes. 4. Discard the supernatant and resuspend the cell pellet in the recommended complete medium (see specific batch information for the appropriate dilution ratio). 5. Incubate the culture under appropriate atmospheric and temperature conditions (see "Culture Conditions" for this cell line). NOTE: It is important to avoid excessive alkalinity of the medium during cell recovery. To minimize this risk, it is recommended to place the culture vessel containing the growth medium in the incubator for at least 15 minutes before adding the vial contents. This allows the medium to stabilize at its normal pH (7.0 to 7.6).
References:	.Shimizu,K., Yamada,M., Moriuchi,S., Tamura,K., Okamoto,Y., Park,KC., Mabuchi,E., Miyao,Y., and Hayakawa,T. Human medulloblastoma cell lines: ONS-76 and ONS-81, Hum. Cell, 3: 265-266, 1990. Tamura,K., Shimizu,K., Yamada,M., Okamoto,Y., Matsui,Y., Park,KC., Mabuchi,E., Moriuchi,S., and Mogami,H. Expression of Major histocompatibility complex on human medulloblastoma cells with neuronal differentiation, Cancer Res., 49: 5380-5384, 1989.
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