

Banco de Células do Rio de Janeiro

Data Sheet

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BCRJ Code: 0321

Cell Line: Reh

Species: Homo sapiens

Vulgar Name: Human

Morphology: Lymphoblast

Disease: Acute Lymphocytic Leukemia (Non-T; Non-B)

Growth Properties: Suspension

Genes Expressed CD3 A (17%) B (17%) C (20%), CD4 (15%), CD10 (55%) Genes Expressed CD3 A (17%) B (17%) C (20%), CD4 (15%), CD10 (55%) Genes **Products:**

Expressed CD3 A (17%) B (17%) C (20%), CD4 (15%), CD10 (55%) Genes

Expressed :CD3 A (17%) B (17%) C (20%), CD4 (15%), CD10 (55%)

Biosafety: 1

RPMI-1640 medium modified to contain 2 mM L-glutamine, 4500 mg/L glucose **Culture Medium:**

and 10% of fetal bovine serum.

Cultures can be maintained by addition or replacement of fresh medium. Start **Subculturing:**

cultures at 2 X 10e5 cells/mL and maintain between 1 X 10e5 and 1 X 10e6

cells/mL.

Subculturing Medium

Renewal:

Every 2 to 3 days

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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:

always be used and a full face mask always be worn when handling frozen vials. It is important to note that some vials leak when submersed in liquid nitrogen and will slowly fill with liquid nitrogen. Upon thawing, the conversion of the liquid nitrogen back to its gas phase may result in the vessel exploding or blowing off its cap with dangerous force creating flying debris. 1. Thaw the vial by gentle agitation in a 37°C water bath. To reduce the possibility of contamination, keep the Oring and cap out of the water. Thawing should be rapid (approximately 2 minutes). 2. Remove the vial from the water bath as soon as the contents are thawed, and decontaminate by dipping in or spraying with 70% ethanol. All of the operations from this point on should be carried out under strict aseptic conditions. 3. For cells that are sensitive to DMSO is recommended that the cryoprotective agent be removed immediately. Transfer the vial contents to a centrifuge tube containing 9.0 mL complete culture medium and spin at approximately 125 x g for 5 to 7 minutes. 4.Discard the supernatant and Resuspend cell pellet with the recommended complete medium (see the specific batch information for the culture recommended dilution ratio). 5. Incubate the culture in a appropriate atmosphere and temperature (see "Culture Conditions" for this cell line). NOTE: It is important to avoid excessive alkalinity of the medium during recovery of the cells. It is suggested that, prior to the addition of the vial contents, the culture vessel containing the growth medium be placed into the incubator for at least 15 minutes to allow the medium to reach its normal pH (7.0 to 7.6).

SAFETY PRECAUTION: Is highly recommend that protective gloves and clothing

References:

reaves M, Janossy G. Patterns of gene expression and the cellular origins of human leukaemias. Biochim. Biophys. Acta 516: 193-230, 1978.

Depositors:

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ATCC:

CRL-8286