

## Protocol for cryopreservation of RPTEC/TERT1

Version: May 2021

Evercyte Ord. No.:	CHT-003-0002
Designation:	RPTEC/TERT1, human renal proximal tubular epithelial cells
Freezing medium:	Cryostor® cell cryopreservation medium CS10 (Sigma-Aldrich, Cat# C2874, ready-to-use)
Additional reagents:	0.05 % Trypsin-EDTA (Gibco, Cat# 25300-054, ready-to-use, stored at 4°C) Defined Trypsin Inhibitor (Gibco, Cat# R007100, ready-to-use, stored at 4°C) PBS (Sigma, Cat# D8537, ready-to-use, stored at RT) ProxUp medium (Evercyte, Cat# MHT-003)
Freezing cells:	<ul style="list-style-type: none"><li>- detach the cells from the culture vessel by using Trypsin-EDTA and Defined Trpsin Inhibitor as described in protocol <i>Passaging of RPTEC/TERT1 cells</i></li><li>- resuspend the detached cells in growth medium and centrifuge at 170 g for 5 min</li><li>- discard the supernatant</li><li>- resuspend the resulting cell pellet in the remaining droplet</li><li>- add freezing medium (tempered to 4°C) to reach a cell density of about <math>1.5-2 \times 10^6</math> cells/ml (for thawing in a 25 cm<sup>2</sup> culture flask)</li><li>- add 1 ml of this cell suspension to each pre-cooled cryovial and immediately transfer the cells to -80°C</li><li>- after 24 hours transfer the vials to the liquid nitrogen tank</li></ul>
Thawing cells:	<p><b>When you start cultivating the cells, please transfer the content of the original Evercyte vial containing RPTEC/TERT1 cells into a T25 roux flask as described in the following:</b></p> <ul style="list-style-type: none"><li>- add 6 ml of growth medium to a 25 cm<sup>2</sup> culture flask and place the culture flask in the incubator for at least 30 min to allow the medium to reach 37°C and its normal pH</li><li>- take a vial of frozen cells, rinse it outside with ethanol and pre-warm in the hand until one last piece of frozen cells is seen</li><li>- then, immediately transfer the content of the vial to a 15 ml centrifugation tube pre-filled with 9 ml of medium pre-cooled to 4°C and centrifuge for 5 min at 170 g</li><li>- discard the supernatant and resuspend the cell pellet in the remaining droplet</li><li>- add 1 ml of the pre-warmed medium to the cells, transfer the cells to the prepared culture flask and incubate at 37°C in a suitable incubator</li><li>- perform a medium change 24 hours after thawing</li><li>- if the cells are already confluent at this point, they have to be passaged as described in protocol <i>Passaging of RPTEC/TERT1 cells</i></li><li>- for the first passages after thawing we recommend a split ratio of 1:2 or lower</li><li>- after thawing, cells generally need 2-3 days before they can be passaged, do not split the cells before having reached about 95% confluence</li></ul>
Related products:	ProxUp ready-to-use medium, 500 ml (Evercyte, Cat# MHT-003) ProxUp basal medium, 500 ml (Evercyte, Cat# MHT-003-B)

ProxUp supplements (Evercyte, Cat# MHT-003-S)