

SW1116 Catalog No: tcel307



Available Sizes

Size: 1×10⁶cells/t25culturebottle



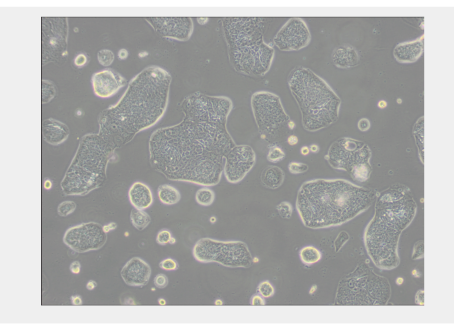
Specifications

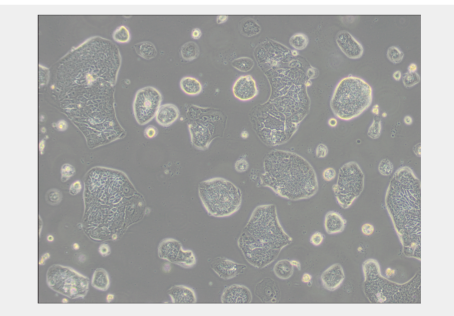
Subculturing:

Cells must be subcultured at about 80% confuency , before they reach 90%. Remove and discard culture medium. Briefly rinse the cell layer with DPBS solution to remove all traces of serum that contains trypsin inhibitor. Add 1.0 to 2.0 mL of Trypsin-EDTA solution to flask and observe cells under an inverted microscope until cell layer is dispersed (usually within 2 to 3 minutes). Cells that are difficult to detach may be placed at 37°C to facilitate dispersal. Add 4.0 to 6.0 mL of complete growth medium and aspirate cells by gently pipetting. Add appropriate aliquots of the cell suspension to new culture vessels.

Product Description

CSAp negative(CSAp-).Colon antigen 3, negative.The cells are positive for keratin by immunoperoxidase staining.The line is positive for expression of c-myc, K-ras, H-ras, myb, sis and fos oncogenes.N-myc and N-ras oncogene expression were not detected.Tumor specific nuclear matrix proteins CC-4, CC-5 and CC-6 are expressed.

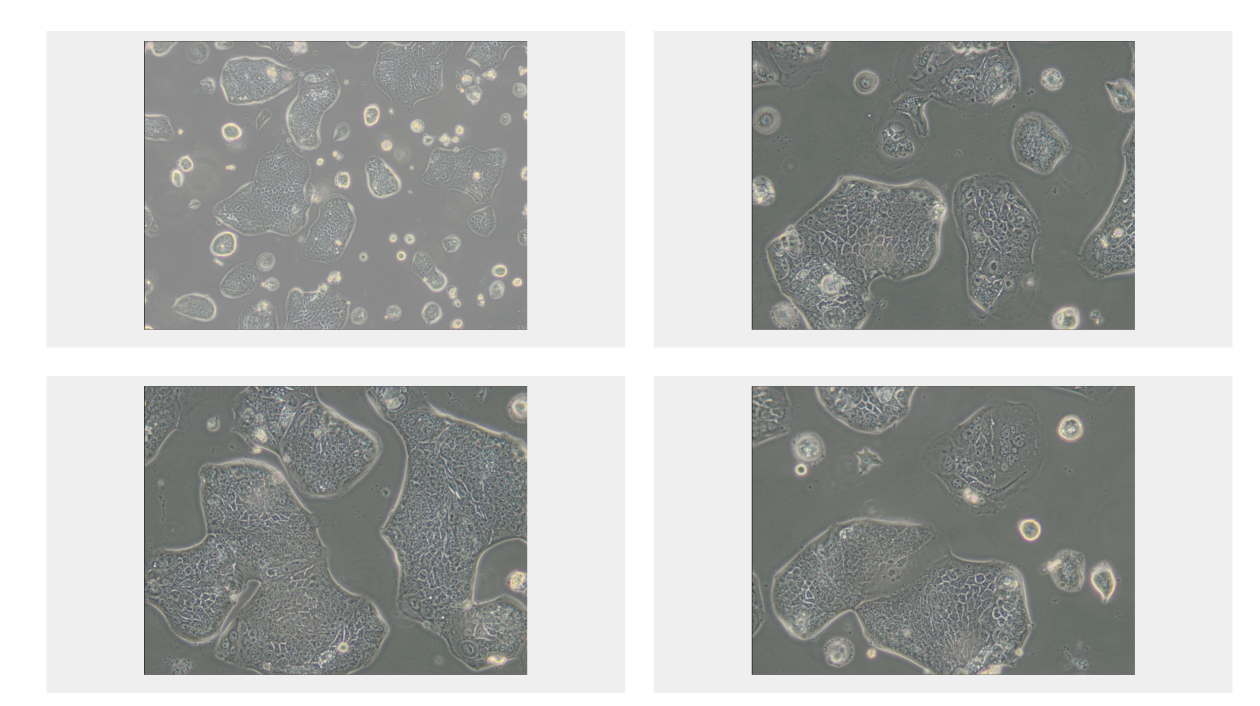




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