

# Adipose-Derived MSC Isolation

This document describes the recommended protocol for isolation of mesenchymal stem cells from subcutaneous adipose tissue, using one of the new CnT-Prime MSC media. For the recommended thawing, passaging, and freezing protocols, please see the General Cultivation Protocol in the resources section of [www.cellntec.com](http://www.cellntec.com).

## General Notes

This protocol describes the isolation of adMSC from subcutaneous adipose tissue, using collagenase 1 (for example Worthington cat#LS004194), and erythrocyte lysis buffer (for example Biologend cat# 420301).

The quantities described in this protocol are for 3 grams of adipose tissue. For isolation from larger amounts, scale up the reagents accordingly.

Recommended culture media for maximum isolation efficiency are #CnT-PR-MS-C or #CnT-PR-MS-C-XF (referred to in the protocol as CnT-Prime MSC Medium)

At the end of the first expansion passage, cell yield is commonly in the range of 3-4 million cells.

## Preparation

Before starting to work with the cells, ensure that the amount of medium, buffers and enzymes required have been pre-warmed at least to room temperature.

Always use sterile instruments, aseptic technique, and work in a laminar flow to maintain sterility.

## Isolation Procedure From Adult Subcutaneous Adipose Tissue

1. For maximum cell yield, store the tissue biopsy in #CnT-XP3 biopsy maintenance medium at 4°C from sampling right up until the start of isolation.
2. Place the adipose tissue into a Petri dish, wash off any blood, and cut out the 3 gram sample using a scalpel.
3. Mince the tissue piece thoroughly with a scalpel, until it is a fully homogenized slurry (5-10 mins).
4. Transfer the homogenized cell slurry into 10 mL of filter sterilized collagenase 1 solution (2 mg/mL), containing 1x antibiotics (#CnT-GAB-5 for maximum heat and pH stability)
5. Incubate for 1.5 hours at 37°C, with gentle vortexing every 30 minutes.
6. Dilute the enzyme solution with 10 mL of CnT-Prime MSC medium, and allow to sit until the fat layer separates. Remove the lower clear solution with a pipette, and filter sequentially through 100 um and 70 um filters.
7. Centrifuge the filtered solution at 200g for 5 minutes, and discard the surface lipid layer and supernatant using a pipette.
8. Resuspend the pellet (stromal vascular fraction) in 1 mL of CnT-Prime MSC medium, then add 3 mL of erythrocyte lysis buffer, and incubate at room temperature for 5-10 minutes (depending on the quantity of RBCs).

9. Add 10 mL of CnT-Prime MSC medium to the lysis buffer, and centrifuge at 200g for 5 minutes. Remove the supernatant, resuspend the cell pellet in 1 mL of medium, and count the cells
10. Seed cells at 20,000 cells per sq cm in CnT-Prime MSC medium with 1x antibiotics, and culture overnight.
11. Perform a medium change the following day, and then every alternate day (Mon/Wed/Fri) until cells reach confluency (typically 6-7 days).
12. Passage the cells just prior to full confluency, re-seeding at 2,000 cells per sq cm. For further culture after passaging it is recommended to omit the antibiotics.

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