

Protocol

Isolation of Progenitor and Stem Cells from Canine Adipose Tissue

- ⚠ All procedures must be performed in biosafety hoods.
- ⚠ This procedure is intended for processing canine subcutaneous adipose tissue only.
- ⚠ Surgically remove the inquinal fat pad using sterile techniques.
- ▲ Using fresh adipose tissue will yield the best results.
- Pre-warm the processing solution (sterile PBS or sterile Lactated Ringer's) to 37°C. Measure the tissue weight in a pre-zeroed sterile container; this will be used to calculate the processing solution volume and Celase® GMP volume.
- Wash the tissue with processing solution if bloody.
- Use surgical scissors to mince the tissue to a 3-5 mm³ size or until no obvious large pieces are observed. Add a small amount of processing solution if the tissue is too dry.
- Reconstitute the Celase GMP vial with 5 mL of the processing solution. Invert the vial several times to fully reconstitute the enzyme into a clear solution (reference Celase GMP IFU).
- Place the washed and minced tissue into a sterile container, usually 6 times the size of the original tissue volume. For a given amount of tissue, add the appropriate volumes of processing solution and Celase GMP to the container; tightly close the container.

| Adipose Tissue | Processing Solution | Celase GMP |
|----------------|---------------------|------------|
| 5 mL | 20 mL | 0.7 mL |
| 10 mL | 40 mL | 1.4 mL |
| 15 mL | 60 mL | 2.1 mL |
| 20 mL | 80 mL | 2.8 mL |
| 25 mL | 100 mL | 3.5 mL |
| 30 mL | 120 mL | 4.2 mL |
| 35 mL | 140 mL | 4.9 mL |
| 40 mL | 160 mL | 5.6 mL |
| 45 mL | 180 mL | 6.3 mL |
| 50 mL | 200 mL | 7.0 mL |

- Place the container in a 37° C shaking water bath for 60 ± 15 minutes until the tissue becomes visibly broken down. If a shaking water bath is not available, gently swirl the container contents by hand continuously or in 5 minute increments for 60 ± 15 minutes.
- Ocllect the non-buoyant solution into multiple 50 mL centrifuge tubes. Place the tubes in a centrifuge and concentrate the cells at 600 g for 5 minutes at room temperature with a low-medium brake speed.
- Carefully aspirate the top lipid, floating layer and the solution that contains Celase GMP.

 ∆ Do not disturb the cell pellet and leave 5-10 mL of the solution in the tube.
- Pesuspend the cell pellet in each 50 mL tube with the remaining solution. Combine the cell pellets from each tube and then wash with the processing solution 2 times to reduce residual enzyme. Resuspend the combined cell pellet with 25-30 mL of the processing solution and pass through a 100 μm strainer followed by a 40 μm strainer.
- Ocentrifuge 1 more time and resuspend with the remaining processing solution into the final desired volume for cell counting and further *in vitro* and *in vivo* studies.

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