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JMN-MSMP Notexin Muscle scRNAseq

ccherry1

¹JMN-MSMP

Cellular Senescence Network (SenNet) Method Development Community

SenNet JMN-MSM



ccherry

ABSTRACT

Murine notexin muscle injury followed by single cell RNA sequencing





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Skeletal muscle injury and collection

1 In a given mouse, both tibialis anterior (TA) muscles were injected with 10 μl of notexin (10 μg/ml; Latoxan, France). 2 At various days post injury, the mouse was sacrificed and the TAs were collected. Each TA was proccessed independently to generate single cell suspensions. **Generate single-cell suspensions** 3 Muscles were enzymatically digested with 8 mg/ml Collagenase D (Roche, Basel, Switzerland) and 10 U/ml Dispase II (Roche, Basel, Switzerland) and then manually dissociated to generate cell suspensions. 4 Myofiber debris was removed by filtering the cell suspensions through a 100 µm and then a 40 µm filter (Corning Cellgro # 431752 and # 431750). 5 After filtration, erythrocytes were removed by incubating the cell suspension in erythrocyte lysis buffer (IBI Scientific #89135-030). 6 After digestion, the single-cell suspensions were washed and resuspended in 0.04% BSA in PBS at a concentration of 10⁶ cells/ml. A hemocytometer was used to manually count the cells to determine the concentration of the suspension. Single-cell RNA library preparation and sequencing 7 Single-cell RNA-sequencing libraries were prepared using the Chromium Single Cell 3' reagent kit v3 (10x Genomics, Pleasanton, CA) following the manufacturer's protocol. 8 Cells were diluted into the Chromium Single Cell A Chip to yield a recovery of 6,000 single-cell transcriptomes with <5% doublet rate. 9 Libraries were sequenced on the NextSeq 500 (Illumina, San Diego, CA).