

Single-nucleus transcriptional and chromatin accessibility analyses of maturing mouse Achilles tendon uncover the molecular landscape of tendon stem/progenitor cells

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eLife Assessment

This study presents a **valuable** finding of novel markers that may potentially identify resident tendon stem/progenitor cells (TSPCs). The study also presents a comprehensive single-cell transcriptional dataset that will be of value to the field. The evidence supporting the identification of novel markers of a TSPC is **incomplete**, requiring clarification of current analyses, additional analyses between ages, and additional validation experiments to demonstrate that these markers are indeed specific and these cells are indeed TSPCs. This work will be of interest to biologists and engineers focused on tendons and ligaments.

<https://doi.org/10.7554/eLife.104768.1.sa3>

Abstract

Tendons and ligaments are crucial connective tissues linking bones and muscles, yet achieving full functional recovery after injury remains challenging. We investigated the characteristics of tendon stem/progenitor cells (TSPCs) by focusing on the declining tendon repair capacity with growth. Using single-cell RNA sequencing on Achilles tendon cells from 2- and 6-week-old mice, we identified *Cd55* and *Cd248* as novel surface antigen markers for TSPCs. Combining single-nucleus ATAC and RNA sequencing analyses revealed that *Cd55* and *Cd248* positive fractions in tendon tissue are TSPCs, with this population decreasing at 1 weeks. We also identified candidate upstream transcription factors regulating these fractions. Functional analyses of isolated CD55/CD248 positive cells demonstrated high clonogenic potential and tendon differentiation capacity, forming functional tendon-like tissue *in vitro*.

This study establishes CD55 and CD248 as novel TSPC surface antigens, potentially advancing tendon regenerative medicine and contributing to the development of new treatment strategies for tendon and ligament injuries.

Introduction

Tendons are connective tissues involved in the contractile movement of muscle to bone. Tendons are rich in extracellular matrix components, such as type 1 collagen and proteoglycans, which both have elastic and viscous properties to withstand overload (Andarawis-Puri et al., 2015; Asahara et al., 2017; Connizzo et al., 2013).

Tendons have a hierarchical structure. Collagen fibrils aggregate to form fibers, which then bundle into fascicles. Each fascicle is surrounded by a thin connective tissue layer called the endotenon. Fascicles are further enclosed by the epitenon, and in some cases, an additional outer layer called the paratenon. The number of fascicles and the presence of a paratenon can vary among different tendons and species (Jozsa et al., 1991; Walia et al., 2019).

Tendon and ligament injuries potentially account for the majority of musculoskeletal disorders and can lead to arthritis and spondylitis (Gracey et al., 2020). The most popular treatment approaches are surgery and conservation (Steinmann et al., 2020). However, full functional recovery is often not achieved due to scarring and fibrosis in the injured area (Guilak et al., 2014; Thomopoulos S et al., 2015). Furthermore, the risk of postoperative rupture is high (Leong et al., 2020; Loiacono et al., 2019). In this context, tendon regeneration focused on tendon stem/progenitor cells (TSPCs) is attracting attention (Leong et al., 2016). Tendons have few cellular components, including tendon cells (tenocytes) and TSPCs (Tang et al., 2016; Bi et al., 2007). Tenocytes are responsible for tendon homeostasis, while TSPCs self-renew and differentiate into tenocytes.

Tendon Stem/Progenitor Cells (TSPCs) were first reported in 2007 by Bi Y et al. (Bi et al., 2007). They demonstrated that TSPCs possess self-renewal capacity, colony-forming ability, and multi-differentiation potential *in vitro*. Furthermore, TSPCs were reported to simultaneously express stem cell markers such as Cd44 and Stem cells antigen-1 (*Ly6a*), as well as tendon-related genes like Scleraxis (*Scx*) and Collagen type I alpha 1 chain (*Col1a1*), and were identified as a subset of tendon cells within the tendon fascicle.

Subsequent research on TSPC localization has yielded diverse perspectives. Harvey et al. reported that *Tppp3/Pdgfra*-positive cells in the epitenon are induced upon tendon injury and contribute to repair (Harvey et al., 2019). Additionally, Yin et al. and Tempfer et al. demonstrated the presence of TSPCs around blood vessels (Yin et al., 2016; Tempfer et al., 2009).

Markers used to characterize TSPCs include CD73, CD105, and CD90, which are known criteria for mesenchymal stem cells (MSCs) (Lui et al., 2015). Additionally, TSPCs have been reported to express POU class 5 homeobox 1 (*Oct-4*), *Nanog*, Nucleostemin, stage-specific embryonic antigen-4 (*SSEA-4*), *c-Myc*, SRY-box transcription factor (*Sox2*), *Fucosyltransferase 4* (*Fut-4*), and other genes (Zhang et al., 2016). Expression of CD146 and CD44 has been confirmed as well (Ruzzini et al., 2014). The *Tppp3/Pdgfra*-positive TSPCs reported by Harvey et al. highly express *Cd34*, which is generally considered to have low expression in MSCs (Harvey et al., 2019; Tachibana, et al., 2022). *Cd34* is known to be highly expressed in mouse embryonic limb buds at E14.5 compared to E11.5 (Havis et al., 2014), making it a potential marker for TSPCs. However, these markers are not specific to TSPCs, thus, a definitive method to distinguish TSPCs from mature tendons *in vivo* has not been established yet (Li et al., 2019; Wang et al., 2008; Chen et al., 2012; Cho et al., 2018; Fang et al., 2022).

Regarding tendon regenerative capacity, Howell et al. reported an interesting observation. Tendons in juvenile mice can regenerate functional tissue after injury, but this ability is lost in mature mice, resulting in scar tissue formation (Howell et al., 2017 [DOI](#)). This finding suggests the possibility of abundant TSPCs in juvenile mouse tendons.

Given this background, we hypothesized that evaluating tendon tissue heterogeneity at the single-cell level using juvenile mouse tendons would enable a more detailed characterization of TSPCs. This approach is expected to advance the identification and characterization of TSPCs, which has been challenging with conventional methods.

Therefore, we performed scRNA-seq using cells collected from 2- and 6-week-old mouse Achilles tendons and investigated clusters that co-express known TSPCs markers, such as *Tppp3*, *Pdgfra*, and *Ly6a*. Then, we analyzed the expression of surface antigens in these clusters and identified *Cd55* and *Cd248* as novel candidate surface antigens for TSPCs.

Furthermore, snATAC-seq and snRNA-seq were performed simultaneously using cells collected from Achilles tendons to evaluate the validity of *Cd55* and *Cd248* as surface antigens for TSPCs and to identify the landscape of transcription factors (TFs) involved in tendon maturation. We sorted mouse Achilles tendon cells based on CD55 and CD248 and confirmed their phenotypes, demonstrating high clonogenicity and highly efficient induction into tendon cells. These results suggested that CD55 and CD248 are novel surface antigens of TSPCs and may be useful for understanding the process of tendon maturation and for applications in cell therapy.

Material and methods

Single cell isolation

Achilles tendons were digested to obtain a single-cell suspension. In total, 40 or 60 (2-week and 6-week, respectively) mouse Achilles tendons were processed together as a single sample. After washing with 1× phosphate-buffered saline (PBS) several times and cutting 1-mm-wide sections using scissors, tendons were digested for 1 h at 37°C with continuous shaking at 1,200 rpm and 37°C in a dissociation solution consisting of 30 mg/mL collagenase (Wako, Osaka, Japan). After digestion, the single-cell suspension was filtered for debris using a 40 µm cell strainer and washed twice with 1× PBS. The samples were centrifuged for 15 min at 1,500 rpm and 4°C. Dead cells were removed using a Dead Cell Removal Kit (Miltenyi Biotec, Bergisch Gladbach, Germany).

scRNA-seq library construction and sequencing

Cells were resuspended in PBS with 1% fetal bovine serum (FBS) at a concentration of 10,000 cells per µL. Then, 10,000 cells per sample were loaded on a Chromium Controller (10x Genomics, Pleasanton, CA, USA) for single cell capture. Libraries were prepared using Single Cell 3' Library & Gel Bead Kit v3 (10x Genomics) following the manufacturer's instructions. A single-cell emulsion (Gel Bead-In-EMulsions, GEMs) was created by making barcoded cDNA unique to each individual emulsion. A recovery agent was added to break GEM and cDNA was then amplified. A library was produced via end repair, dA-tailing, adapter ligation, post-ligation cleanup with SPRIselect, and sample index PCR. The quality and concentration of the amplified cDNA were evaluated using the Bioanalyzer (Agilent 2100) on a High Sensitivity DNA chip (Agilent, #5065-4401; Santa Clara, CA, USA). Sequencing was performed using the HiSeq X system (Illumina Inc., San Diego, CA, USA) to generate 28/90 bp paired-end reads.

Cell dissociation, nuclei isolation, and snRNA-seq and snATAC-seq library construction and sequencing

After obtaining cells from the mouse Achilles tendons, nuclei were isolated following the Chromium Next GEM Single Cell Multiome ATAC + Gene Expression Reagent Bundle (10XGenomics) and all of the buffers were made according to the manufacturer's instructions. Briefly, after centrifugation, the supernatant was discarded, and the cells were re-suspended in 1 mL of 1% FBS in PBS. Approximately 500,000 cells were transferred to a new tube for further lysis. To remove the supernatant, cells were centrifuged again for 5 min at 300 rcf and 4°C and resuspended in 100 mL of chilled lysis buffer. Cells were lysed for 9 min on ice and 1 mL of wash buffer was added to stop the reaction. Then, the suspension was filtered through a 40 µm cell strainer, cells were centrifuged and resuspended in 66.2 mL of chilled Diluted Nuclei Buffer aiming to target 7000 nuclei. The final concentration of nuclei was determined, followed by transposition, GEM generation, barcoding, and library construction according to the Chromium Next GEM Single Cell Multiome ATAC + Gene Expression (10XGenomics). Libraries were sequenced with the parameters recommended by the manufacturer, using the NovaSeq 6000 (Illumina Inc.) to generate 28/90 bp paired-end reads for gene expression and 50/49 bp paired-end reads for ATAC sequencing.

scRNA-seq and snRNA-seq analyses

Sequencing reads were processed with the Cellranger_arc (10X Genomics, v2.0.0) using the mouse reference mm10. From the gene expression matrix, downstream analyses were carried out using R. Quality control, filtering, data clustering and visualization, and a differential expression analysis were carried out using Seurat (Butler et al., 2018 [DOI](#)). For each dataset, cells with unique feature counts over 2,500 or less than 200 and >5% mitochondrial counts were filtered. Then, heterotypic doublets (assuming 5% of barcodes represent doublets) were removed using DoubletFinder (McGinnis et al., 2019 [DOI](#)).

Unsupervised shared nearest neighbor (SNN) clustering was performed with varying resolution and the results were visualized using uniform manifold approximation and projection (UMAP) (Becht et al., 2019 [DOI](#)). Differentially expressed genes (DEGs) among each cell cluster were determined using the FindAllMarkers function in Seurat. DEGs expressed in at least 25% of cells within the cluster and with a log(fold change) of more than 0.25 were identified as marker genes for the cluster.

Pseudotime analysis

Monocle3 (Trapnell et al., 2014 [DOI](#)) was used to convert the snRNA-seq dataset into a cell dataset object (CDS), preprocess data, correct for batch effects, embed with dimensional reduction, and perform pseudotemporal ordering. Cicero (Pliner et al., 2018 [DOI](#)) was used to generate pseudotemporal trajectories for the snATAC-seq dataset.

Cell-cell communication analysis

Intercellular communication networks were quantitatively inferred and analyzed using scRNA-seq data. The R package CellChat (Jin et al., 2021) was used to visualize the interactions among different cell groups. Two hundred twenty-nine signaling pathway families were grouped as a library to analyze cell-cell communication.

GO analysis

The R package ClusterProfiler (Yu et al., 2012 [DOI](#)) was used to perform a gene enrichment analysis. The p-values were corrected by the Benjamini & Hochberg method.

Motif enrichment analysis (ChIP seeker)

Genomic regions containing snATAC-seq peaks were annotated using ChIPSeeker (Yu et al., 2015) and the UCSC database on mouse (mm10).

snATAC-seq analysis

The Cell Ranger ATAC pipeline (1.2.0) (Satpathy et al., 2019) was used to preprocess the data resulting from sequencing. First, Tn5 sites were mapped to the mouse reference transcriptome mm10, and duplicate reads and background cells were removed. This returned barcoded fragment files, which were loaded into Signac (Stuart et al., 2021) for downstream analyses using the standard Signac/Seurat pipeline. Macs2 (Zhang et al., 2008) was run on the fragment files to call peaks using the Signac ‘CallPeaks’ function. Fragments were mapped to the Macs2 called peaks and assigned to cells using the Signac ‘FeatureMatrix’ function. Nucleosome signal strength and transcription start site (TSS) enrichment for each cell were calculated using the Signac ‘NucleosomeSignal’ and ‘TSSEnrichment’ functions, respectively. Outliers in the QC metric categories were removed as per Signac’s standard processing guidelines. Latent semantic indexing (LSI), a form of dimensional reduction, was performed using the Signac ‘RunTFIDF’ and ‘RunSVD’ functions. The UMAP hyperparameters were varied to produce consistent object shapes (using R). Once hyperparameters were chosen, the Signac/Seurat’s ‘RunUMAP’ function was run on the LSI dimensions chosen earlier for UMAP embedding. The Signac/Seurat ‘FindNeighbors’ function was run using the same LSI dimensions used for UMAP to compute the nearest neighbor graph. Signac/Seurat ‘FindClusters’ was then run at varying resolutions. A gene activity matrix was constructed by counting ATAC peaks within the gene body and 2 kb upstream of the transcriptional start site for protein-coding genes annotated in the Ensembl database. The gene activity matrix was log-normalized prior to label transfer with the aggregated snRNA-seq Seurat object using a canonical correlation analysis. The aggregated snATAC-seq object was filtered using a 97% confidence threshold for cell-type assignment following label transfer to remove heterotypic doublets. The filtered snATAC-seq object was reprocessed with TFIDF, SVD, and batch effect correction followed by clustering and annotation based on lineage-specific gene activity. Differential chromatin accessibility between cell types was assessed with the Signac FindMarkers function for peaks detected in at least 20% of cells using a likelihood ratio test and a log-fold-change threshold of 0.25. Bonferroni-adjusted p-values were used to determine significance at an FDR < 0.05.

Integration of snRNA-seq and snATAC-seq data

snRNA-seq and snATAC-seq data were integrated using the cluster-label transfer procedure as implemented in Signac and Seurat. Each snRNA-seq sample, as clustered individually and its cluster labels were projected onto the matching, individually clustered snATAC-seq sample or vice versa. Anchors were identified for condition-matched snRNA- and snATAC-seq samples using the FindTransferAnchors function and a canonical correlation analysis (CCA) was performed using the snRNA expression values and the snATAC-imputed gene expression values. The anchors were used to transfer cluster-label identifiers between the two data types using the TransferData function. Each cell in the query was assigned the cluster label with the highest prediction score, and label transfer was considered successful for query cells with prediction scores above 0.3.

SCENIC analysis

To identify TFs and characterize cell states, a cis-regulatory analysis was performed using the R package SCENIC (Lake et al., 2018), which infers gene regulatory networks based on co-expression and DNA motif analyses. The network activity was then analyzed for each cell to identify recurrent cellular states. In short, TFs were identified using GENIE3 and compiled into modules

(regulons), which were subsequently subjected to cis-regulatory motif analysis using RcisTarget with two gene-motif rankings: 10 kb around the TSS and 500 bp upstream. Regulon activity in every cell was then scored using AUCell.

Immunohistochemistry

Tissue samples were fixed in 4% paraformaldehyde overnight at 4°C, washed in 1× PBS, cryo-preserved in 20% sucrose overnight, and embedded in optimal cutting temperature (OCT) compound (4583; Sakura Finetek Japan Co., Ltd., Tokyo, Japan). Then, the bio-tendon was cryo-sectioned at a thickness of 10 µm and desiccated by air-drying overnight. Immunohistochemical staining was performed using a Vectastain ABC-AP Rabbit IgG Kit (AK-5001; VECTOR LABORATORIES, INC., Burlingame, CA, USA) and Vector Red (SK-5100; VECTOR LABORATORIES, INC.) according to the manufacturer's instructions. Anti-CD55 (1/500 dilution) (ab231061; Abcam, Cambridge, UK) and Anti-CD248 antibodies (1/500 dilution) (LS-B2712; LSBio, Seattle, WA, USA) were used as the primary antibodies.

Fluorescence-activated cell sorting (FACS)

Cells harvested from the Achilles tendon were incubated for 60 min on ice with APC-CD55 (#131812; BioLegend, San Diego, CA, USA) and a primary antibody against CD248 (#LS-B2712) using a FACS buffer (PBS containing 1 [v/v%] FBS). After washing, cells were reacted with fluorescent-conjugated rabbit secondary antibody. Cell sorting was performed using MoFlo XDP FACS (Beckman Coulter, Brea, CA, USA).

Primary cell culture

Primary TSPCs were isolated from the Achilles tendons of 2-week-old mice with collagenase digestion described above. Single-cell suspensions were cultured in the culture medium (MEMα + 20% FBS + 1% penicillin-streptomycin + 1 [v/v%] 100× non-essential amino acid solution [NEAA; Gibco], 1 [v/v%] 100× GlutaMAX [Gibco]). At 80%–90% confluence, cells were trypsinized, centrifuged, resuspended in culture medium as passage 1 cells, and incubated in 5% CO₂ at 37°C, with fresh medium every 2–3 days.

Colony formation assay

For the colony formation assay, single-cell suspensions of TSPCs (1000 cells/well) were seeded and incubated in six-well plates for 12–14 days in the growth medium and fixed with 4% paraformaldehyde (PFA) (Sigma-Aldrich, St. Louis, MO, USA). Then, 0.1% crystal violet solution (Wako) was used to stain the cells. Colonies of >30–50 cells were defined as a single colony unit, and the number of clusters was counted using the ImageJ package ColonyArea (Guzmán et al., 2014 [\[4\]](#))

RT-PCR

Total RNA was extracted using TRIzol reagent (Invitrogen, Grand Island, NY, USA). The PrimeScript RT Reagent Kit (Takara, Tokyo, Japan) was used for reverse transcription of mRNA. Complementary DNA was quantitated by qRT-PCR using a Thunderbird SYBR mix (Toyobo Co., Osaka, Japan). *B2MG* expression served as a control and changes in gene expression were quantified using the $\Delta\Delta CT$ method.

Tenocytes, cartilage, and osteocyte differentiation

For the differentiation experiment, TSPCs were cultured in 6-well plates (50,000 cells/well). Osteogenic, chondrogenic, and tenogenic differentiation were induced using a corresponding differentiation medium. The osteogenic differentiation medium contained a culture medium supplemented with 10 nM dexamethasone (Sigma-Aldrich), 5 mM β-glycerophosphate (APEXBIO), and 0.05 mM L-ascorbic acid 2-phosphate (Sigma-Aldrich). The chondrogenic differentiation

medium contained a culture medium supplemented with 100 nM dexamethasone and 10 ng/mL BMP2 (Sigma-Aldrich). The tenogenic differentiation medium contained a culture medium supplemented with 10 ng/mL TGF- β 1 (Peprotech, Rocky Hill, NJ, USA), 10 ng/mL GDF-5 (R&D Systems, Minneapolis, MN, USA), and 0.05 mM L-ascorbic acid 2-phosphate. After a 2 week induction period, cells were harvested for RT-PCR.

Bio-cultured tendon (Bio-tendon)

Cultured cells were embedded in a 3D-culture cocktail (Tsutsumi et al., 2022 [DOI](#)). The 3D-culture cocktail was constructed by mixing collagen gel [final concentrations: 2 mg/mL Cellmatrix (Type I-A, Nitta Gelatin Inc., Osaka, Japan) and 1 $\times$ collagen neutralization buffer (Type I-A, Nitta Gelatin Inc.)], pro-survival cocktail (final concentrations: 100 nM B-cell lymphoma extra-large (Bcl-XL) BH4 4-23 (197217-1MG, Calbiochem, United States), 100 μ M carbobenzoxy-valyl-alanyl-aspartyl-[O-methyl]-fluoromethylketone (Z-VAD-FMK) (Promega, Madison, WI, USA)), and culture medium. The 3D chamber was coated with 1% gelatin and incubated at 37°C and 5% CO₂ for 30 min. After washing thrice with 1 $\times$ PBS, 1.0×10^6 cells were transferred to a 3D-culture cocktail mixture and incubated at 37°C and 5% CO₂ for 60 min for gelation. Following gelation, tenogenic differentiation medium was added to the chamber. Following 24 h of incubation, the 3D-cultured samples were set into a mechanical cell stretch system device (Shellpa Pro, Menicon Co., Ltd./Life Science Department, Aichi, Japan). Cyclic mechanical stretch was performed for 1 week by gradually increasing the stretch loading rate as follows: 1% (day 1), 2% (day 2), 3% (day 3), 4% (day 4), and 5% (days 5–7). The cyclic mechanical stretch was programed at 0.25 Hz for 18 h/day, followed by resting for 6 h/day at 37°C and 5% CO₂. Daily medium changes were also required.

Decellularization by HHP and chemical treatment

The cultured bio-tendon was placed into a plastic pack filled with saline and sealed to prevent implosion and leakage during the procedure. The pack was then pressurized at 1,000 MPa at 30°C for 10 min using an HHP machine (Dr. CHEF; Kobe Steel, Ltd., Hyogo, Japan). After pressurization, the bio-tendon was washed thrice with 30% ethanol (EtOH) by continuous shaking for 5 min at each step. Finally, 1-ethyl-3-(3-dimethylaminopropyl) carbodiimide (EDC)/N-hydroxysuccinimide (NHS)-based cross-linking was performed by adding 70 mM EDC (Wako) and 70 mM NHS (Wako) with 30% EtOH for 24 h at 4°C. The processed bio-tendon was incubated in 1 $\times$ PBS at 4°C until the experiment.

Electron microscopy

Tendon/ligament-like tissues were dissected and fixed in 2.5% glutaraldehyde in 0.1 M phosphate buffer (PB) overnight. For transmission electron microscopy (TEM), the specimens ($n = 3$) were washed with 0.1 M PB, post-fixed in 1% osmium buffered with 0.1 M PB for 2 h, and dehydrated in a graded series of ethanol. Then, the specimens were embedded in Epon 812, sliced into ultrathin sections (70 nm), collected on copper grids, and double-stained with uranyl acetate and lead citrate. The specimens were observed using TEM (JEM-1400Flash; JEOL, Ltd., Tokyo, Japan). For scanning electron microscopy (SEM), the specimens ($n = 2$) were dried in a critical-point drying apparatus (HCP-2; Hitachi, Ltd.) with liquid CO₂ and were sputter-coated with platinum. Then, the specimens were observed using SEM (JSM-7900F/JED-2300; JEOL, Ltd.). Fiber alignment in the microscopic sections was analyze using the OrientationJ image processing tool (an ImageJ plugin).

Stretch test

The mechanical properties were measured using a creep meter (RE-3305S; Yamaden, Tokyo, Japan). After measuring the initial length (mm), diameter (mm), and thickness (mm) using a micrometer, the samples were fixed with two grips, which were pulled at a constant speed of 0.05 mm/s until failure, and the tensile strength (N) and failure strain (mm) were measured. The cross-sectional area (mm²) was calculated using the initial diameter and thickness. The stiffness was manually determined from the slope in the linear region of the failure-stress curve.

The tensile strength (MPa), failure strain (%), stiffness, and Young's modulus were calculated using the following formulas:

Tensile strength (MPa) = tensile strength (N)/cross-sectional area (mm²).

Failure strain (%) = failure strain (mm)/initial length (mm).

Stiffness = Stress (N)/Strain (mm)

Young's modulus = stiffness × initial length (mm)/cross-sectional area (mm²)

Statistical analysis

All values are presented as means ± SEM. Statistically significant differences were assessed using unpaired two-tailed Student's *t*-tests and one-way analysis of variance (ANOVA) with Tukey's post hoc tests. Statistical significance was set at *p* < 0.05.

Results

Identification of Novel Surface Antigens for Tendon Stem/Progenitor Cells (TSPCs)

In mice, the healing capacity for tendon injuries is high up to 2 weeks of age, but decreases as the musculoskeletal system matures at 6 weeks (Howell et al., 2017 [↗](#)). To investigate whether this phenomenon is due to changes in the cellular population of tendon tissue, including fluctuations in progenitor cells, we performed single-cell RNA sequencing on mouse Achilles tendons at these two time points.

Achilles tendons were harvested from mice at 2 and 6 weeks of age. Following collagenase digestion and dead cell removal, we employed massively parallel, droplet-enabled scRNA-seq analysis (10X Genomics Chromium). Data processing was conducted using the Cell Ranger pipeline (10X Genomics). We analyzed 10,314 cells from 2-week-old mice (median 3,167 genes/cell and 41,611 mean reads/cell) and 6,513 cells from 6-week-old mice (median 732 genes/cell and 60,386 mean reads/cell). After doublet removal using Doubletfinder, we merged the datasets and performed unbiased clustering using Seurat, identifying a total of 15 clusters (**Fig. 1A** [↗](#)). The top differentially expressed genes (DEGs) for each cluster, based on log2 fold change and statistical significance, are summarized in **Fig. 1B** [↗](#).

We identified two clusters (0 and 11) with high expression of *Mkx* and *Scx*, transcription factors involved in tendon development and growth. Cluster 0 showed higher expression of ECM-related genes such as *Col1a1* and *Fmod* compared to cluster 11, suggesting these were tenocytes (TC). Cluster 11, with high Periostin (*Postn*) expression, was classified as myotendinous junction (MTJ). Other clusters were characterized as follows: cartilage (CA, high SRY-box transcription factor 9 (*Sox9*) expression), lymphocytes (LC1, LC2, high Lysozyme 2 (*Lyz2*) and Complement component 1, q subcomponent, alpha polypeptide (*C1qa*)), endothelial cells (EC, high Platelet and endothelial cell adhesion molecule 1 (*Pecam1*)), red blood cells (RBC, high Hemoglobin alpha, adult chain 1 (*Hba-a1*)), smooth muscle cells (SM, high Myosin light chain 9 (*Myl9*)), proliferating cells (PC, high *Mki67* and *Stathmin 1* (*Stmn1*)), Schwann cells (SW, high Myelin basic protein (*Mbp*)), vascular endothelial cells (VEC, high Multimerin 1 (*Mmrn1*)), and macrophages (MC, high Protein tyrosine phosphatase receptor type C (*Ptprc*)). In cluster3, *Col1a1* and *Fmod* were expressed; however, *Gm42418* and *AY036118* were highly expressed. These long non-coding RNAs are related to RN45s and therefore cluster3 represents ribosomal contamination (Isola et al., 2024 [↗](#)).

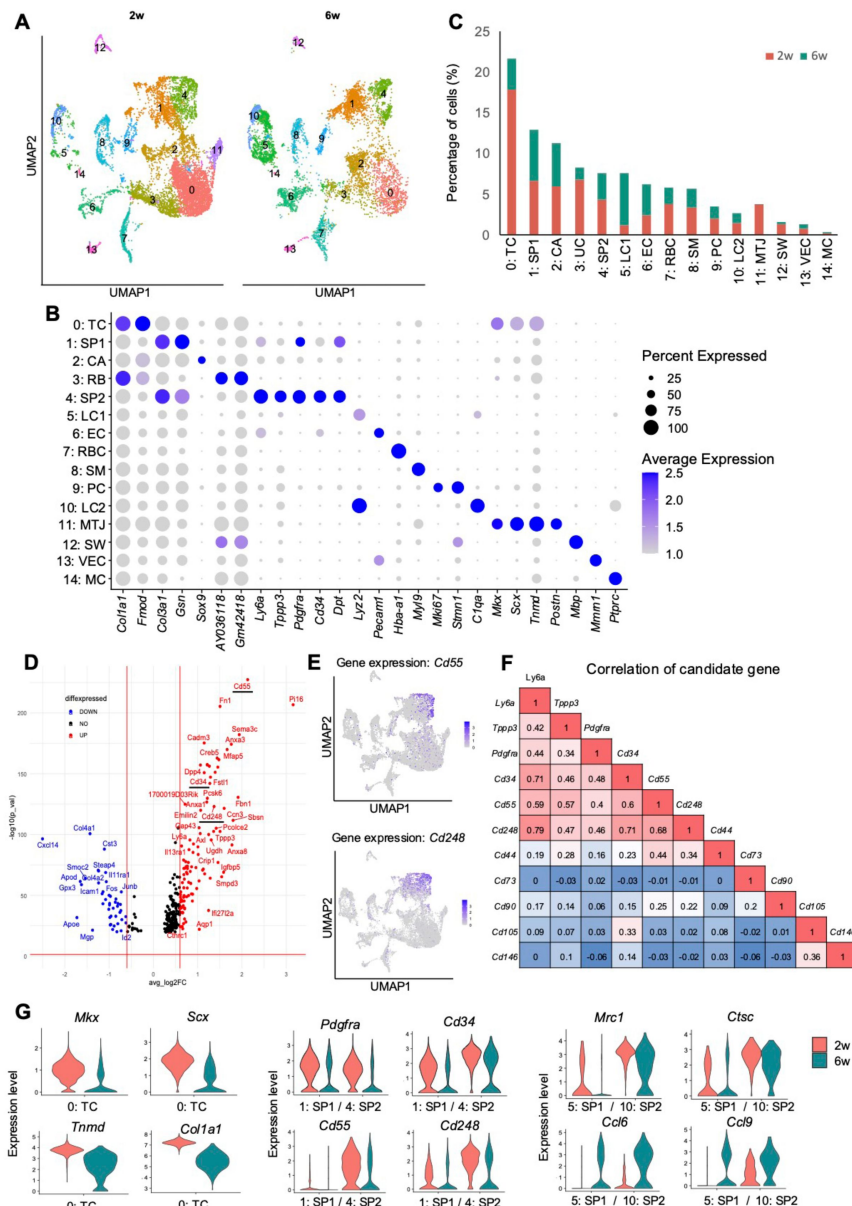


Figure 1.

scRNA-seq of tendon cells from 2-week-old and 6-week-old mice and the identification of surface markers of TSPC.

(A) Integrated uniform manifold approximation and projection (UMAP) scRNA-seq clustering of cells harvested from 2-week-old and 6-week-old mouse Achilles tendons.

(B) Dot plot of average gene expression levels of the indicated genes in each scRNA-seq cluster. The size of the dot reflects the percentage of cells in the cluster that express each gene. TC, tenocyte; SP1, tendon stem/progenitor cell_1; CA, cartilage; RB, ribosomal RNA; SP2, tendon stem/progenitor cell_2; LC1, lymphocyte_1; EC, endothelial cell; RBC, red blood cell; SM, smooth muscle cell; PC, proliferating cell; LC2, lymphocytes_2; MTJ, myotendinous junction cell; SW, Schwann cell; VEC, vascular endothelial cell; MC, macrophage.

(C) Proportions of cells in clusters identified from scRNA-seq. Clusters are colored according to cluster type.

(D) Volcano plot of gene expression in the SP2 cluster and the identification of candidate TSPC marker genes (red under line).

(E) Feature plot of Cd55 and Cd248 expression.

(F) Correlation of gene expression of TSPC candidate genes in 2-week data.

(G) Violin plots presenting the gene expression changes for a selection of differentially expressed genes.

To identify progenitor populations within these clusters, we analyzed expression patterns of previously reported markers *Tppp3* and *Pdgfra* (Harvey et al., 2019 [DOI](#); Tachibana, et al., 2022 [DOI](#)), along with the known stem/progenitor cell marker *Ly6a* (Holmes et al., 2007; Sung et al., 2008 [DOI](#); Hittinger et al., 2013 [DOI](#); Sidney et al., 2014 [DOI](#); Fang et al., 2022 [DOI](#)). We identified subclusters within clusters 1 and 4 showing high expression of these genes, which we defined as SP1 and SP2. SP2 exhibited the highest expression of these genes, suggesting it had the strongest progenitor characteristics.

SP2 also showed strong expression of genes associated with early tendon development in mouse embryos, as identified by RNA-seq of mouse limb tendon cells at E14.5 (Havis et al., 2014 [DOI](#)) and the Eurexpress mouse embryo transcriptome atlas database (Diez-Roux et al., 2011 [DOI](#)) (**Supplemental Figure 1A** [DOI](#)). Gene Ontology analysis revealed upregulation of tendon development-related pathways in SP2, including TGF β production and collagen-containing extracellular matrix (**Supplemental Figure 1B** [DOI](#)). Evaluation of cell numbers in each cluster showed a decrease in SP2, TC, and MTJ at 6 weeks (**Fig. 1C** [DOI](#)).

We then examined DEGs in SP2 compared to SP1 (**Fig. 1D** [DOI](#)). Among these, we identified *Cd34*, *Cd55*, and *Cd248* as genes encoding surface antigens. While *Cd34* has been previously reported in tendon stem progenitor populations (Harvey et al., 2019 [DOI](#); Havis et al., 2014 [DOI](#)), to our knowledge, *Cd55* and *Cd248* have not been associated with this context. Expression of *Cd55* and *Cd248* was localized to areas corresponding to SP1 and SP2 (**Fig. 1E** [DOI](#)). We evaluated the correlation coefficients between the expression of these genes and previously suggested TSPC markers (*Cd73*, *Cd90*, *Cd105*, *Cd44*, *Cd146*) as well as *Tppp3*, *Pdgfra*, and *Ly6a* at 2 weeks of age. *Cd34*, *Cd55*, and *Cd248* showed high correlation, suggesting these genes as new candidate surface antigens for TSPCs (**Fig. 1F** [DOI](#)).

Comparing gene expression between 2 and 6 weeks, we observed a decrease in *Mkx* and *Scx* expression, consistent with previous reports (Grinstein et al., 2019 [DOI](#)). Furthermore, the expression of our candidate tendon progenitor cell markers *Cd34*, *Cd55*, and *Cd248* all decreased at 6 weeks. Inflammatory cells, including macrophages, increased at 6 weeks (**Fig. 1G** [DOI](#)). Gene expression analysis revealed a shift from M2 macrophage-related genes (Mannose receptor C-type 1 (*Mrc1*) and Cathepsin C (*Ctsc*)) at 2 weeks to M1 macrophage-related genes (Chemokine (C-C motif) ligand 6 (*Ccl6*) and *Chemokine (C-C motif) ligand 9* (*Ccl9*)) at 6 weeks. Analysis of cell-cell communication changes in postnatal tendon and surrounding tissues (Lui et al., 2019) using CellChat (Jin et al., 2021) showed a significant decrease in overall interaction at 6 weeks compared to 2 weeks. These results demonstrate that the cellular profile of tendon tissue changes dramatically at 6 weeks, with decreased expression of stem progenitor markers and reduced interaction with surrounding tissues.

Single Nucleus RNA + ATAC Analysis of 2-Week-Old Mouse Achilles Tendon Cells

Gene regulation is mediated by the binding of transcription factors to cis-regulatory elements proximal to the gene. Consequently, epigenetic changes such as chromatin accessibility play a crucial role in gene expression (Miller et al., 2013). Moreover, transcription factors are typically expressed at low levels, potentially leading to false negatives in scRNA-seq due to detection limits. Chromatin accessibility changes often precede gene expression changes, potentially allowing us to predict transcriptional changes (Ranzoni et al., 2020). Therefore, we performed simultaneous ATAC-seq and RNA-seq at the single-cell level to identify gene expression changes and their associated epigenetic alterations, aiming to elucidate the post-natal growth mechanisms of tendon tissue and evaluate the validity of *Cd55* and *Cd248* as markers.

We extracted nuclei from cells isolated from 2-week-old mouse Achilles tendons and conducted droplet-enabled multi-omics analysis (10X Genomics Chromium), performing simultaneous snATAC-seq and snRNA-seq as a reference. We first analyzed the snATAC-seq data. After mapping reads to the genome and calling peaks, we annotated the location of peaks in terms of genomic features. The peaks were associated with promoters, introns, exons, and intergenic regions.

We evaluated 6,571 cells (median high-quality fragments per cell: 17,408) and clustered them using latent semantic indexing and Uniform Manifold Approximation and Projection (UMAP) with the R package Signac (Stuart et al., 2021 [DOI](#)) (**Figure 2A** [DOI](#)). Given the limited knowledge of cell-specific chromatin accessibility, we assessed gene activity by computing counts per cell within the gene body and promoter of protein-coding genes. Using this gene activity data, we performed unsupervised clustering and identified 17 clusters (ground-truth annotation). Clusters expressing tendon-related genes such as *Mkx* and *Scx* were identified as clusters 1 and 6, defined as A1: TC1 and A6: TC2, respectively. Clusters 0, 14, and 15 were identified as expressing stem progenitor markers including *Cd34*, *Ly6a*, *Pdgfra*, *Cd55*, and *Cd248*, and were defined as A2-0: SP1, A2-14: SP2, and A2-15: SP3, respectively. The characteristic gene activities and annotations for each cluster are summarized in **Fig. 2B** [DOI](#).

Next, we analyzed the snRNA-seq data. We evaluated 10,314 cells (median 3,167 genes/cell and 41,611 mean reads/cell). After doublet removal, we performed unbiased clustering using Seurat and identified 17 clusters (**Supplemental Figure 2A** [DOI](#)). Clusters expressing tendon-related genes *Mkx* and *Scx* were clusters 1 and 13, defined as R2-1: TC_1 and R2-13: TC_2, respectively. Clusters 2, 3, and 7 were identified as expressing stem/progenitor-related genes and defined as R2-2: SP_1, R2-3: SP_2, and R2-7: SP_3, respectively. Using the Signac package's "FindTransferAnchors" function, we calculated predicted IDs for snATAC-seq clusters based on snRNA-seq annotations (predicted annotation) (**Fig. 2C** [DOI](#)). Evaluation of the two annotation methods for snATAC-seq revealed that both methods allowed annotation of major cell types, and correlation between the two was maintained. Thus, the validity of predicted annotations based on snRNA-seq was confirmed (**Figure 2D** [DOI](#)).

R2-7: SP3, considered the most undifferentiated stem/progenitor fraction in snRNA-seq, could be divided into A2-0: SP1 and A2-14: SP2 in snATAC-seq. Comparing gene activity of tendon-related and stem/progenitor genes in each cluster of the ground truth annotation, A2-14: SP2 showed high expression of *Ly6a*, *Pdgfra*, and *Tppp3*, and inverse correlation with *Mkx*, *Scx*, and *Tnmd*. Furthermore, the newly identified surface antigens *Cd55* and *Cd248* also showed high expression in cluster 14. In contrast, *Cd34* did not show a high expression pattern in A2-14: SP2. These results suggest that A2-14: SP2 is the most immature cluster, with *Cd55* and *Cd248* showing characteristic expression (**Fig. 2E** [DOI](#)). Subcluster analysis of R7, considered the most immature fraction in snRNA-seq, revealed two clusters based on *Cd55/Cd248* and *Cd34* expression, similar to the snATAC-seq analysis. The former (R7-1) showed high expression of *Ly6a*, *Tppp3*, and *Pdgfra* (**Fig. 2F** [DOI](#)). Trajectory analysis using R2-7 and A2-14 as roots in snRNA-seq and snATAC-seq, respectively, showed increased expression of tendon-related genes *Mkx* and *Scx* along the trajectory, while expression of *Tppp3*, *Cd55*, and *Cd248* decreased (**Figure 3A** [DOI](#)). Evaluation of peaks in each cluster based on gene expression levels in snRNA-seq revealed multiple peaks with heights correlated to gene expression for *Mkx* and *Scx*. Similar evaluation of *Cd55* and *Cd248* showed multiple peaks upstream of the TSS for *Cd248* (**Fig. 3B** [DOI](#)). These results indicate correlation between snRNA-seq and snATAC-seq, and consistent with scRNA-seq results, suggest that A2-14 represents the most undifferentiated TSPCs based on *Cd55* and *Cd248* expression.

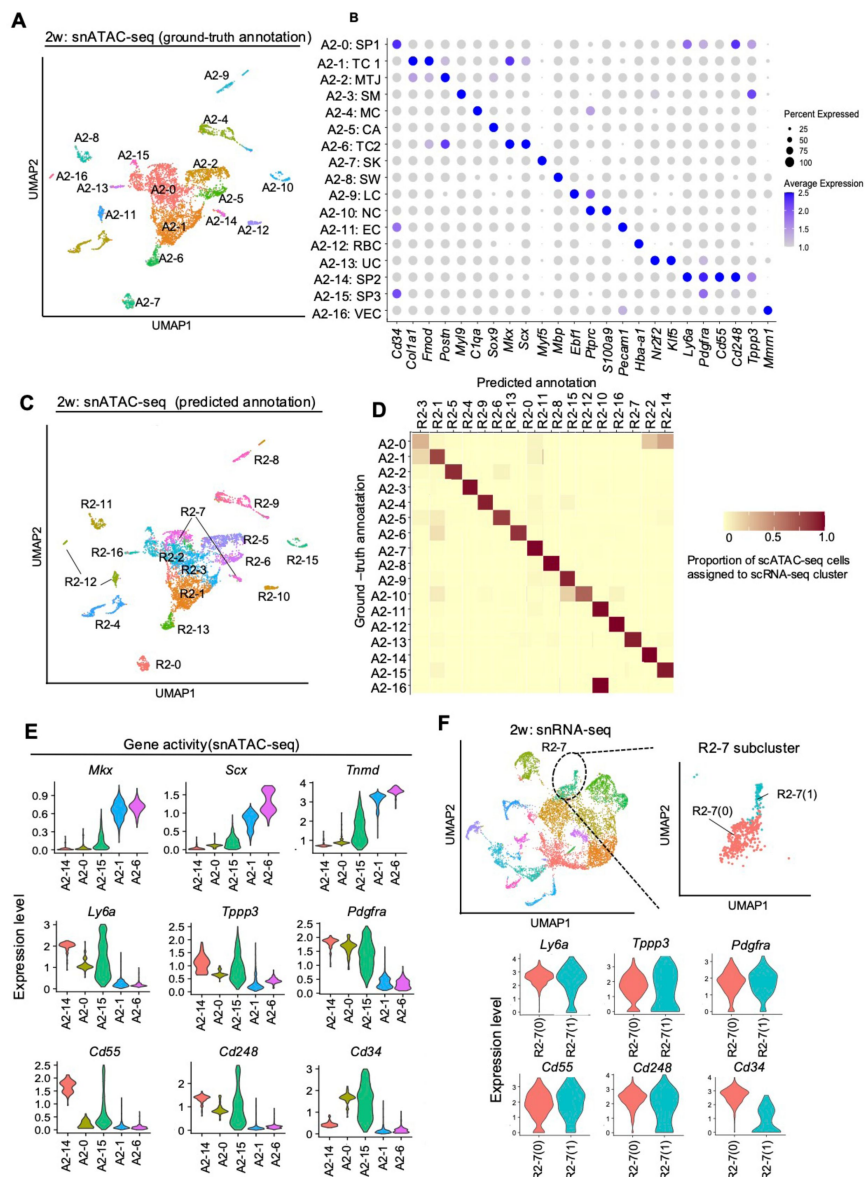


Figure 2.

snATAC-seq of tendon cells from a 2-week-old mouse and the validation of Cd55 and Cd248 as candidate markers of TSPC.

(A) UMAP snATAC-seq clustering of cells derived from the Achilles tendon of a 2-week-old mouse. Annotation was based on each gene activity (ground-truth annotation). SP1, tendon stem/progenitor cell₁; TC1, tenocyte₁; MTJ, myotendinous junction cell; SM, smooth muscle cell; MC, macrophage; CA, cartilage; TC2, tenocyte₂; SK, skeletal muscle cell; SW, Schwann cell; LC, lymphocyte; NC, neutrophil; EC, endothelial cell; RBC, red blood cell; UC, unspecified cell; SP2, tendon stem/progenitor cell₂; SP3, tendon stem/progenitor cell₃; VEC, vascular endothelial cell.

(B) Dot plot of average gene activity of the indicated genes in each snATAC-seq cluster. The size of the dot reflects the percentage of cells in the cluster that express each gene.

(C) UMAP visualization and predicted annotation of 2-week snATAC-seq after integration and label transfer of 2-week snRNA-seq data.

(D) Identification of matching cell clusters between the 2-week snRNA- and 2-week snATAC-seq data from visualized as heatmap. The heatmap shows the proportions of cells from each snATAC-seq cluster across all sample conditions assigned to each snRNA-seq cluster as part of the label-transfer process.

(E) Violin plot of tenocytes and TSPC-related gene expression in each cluster.

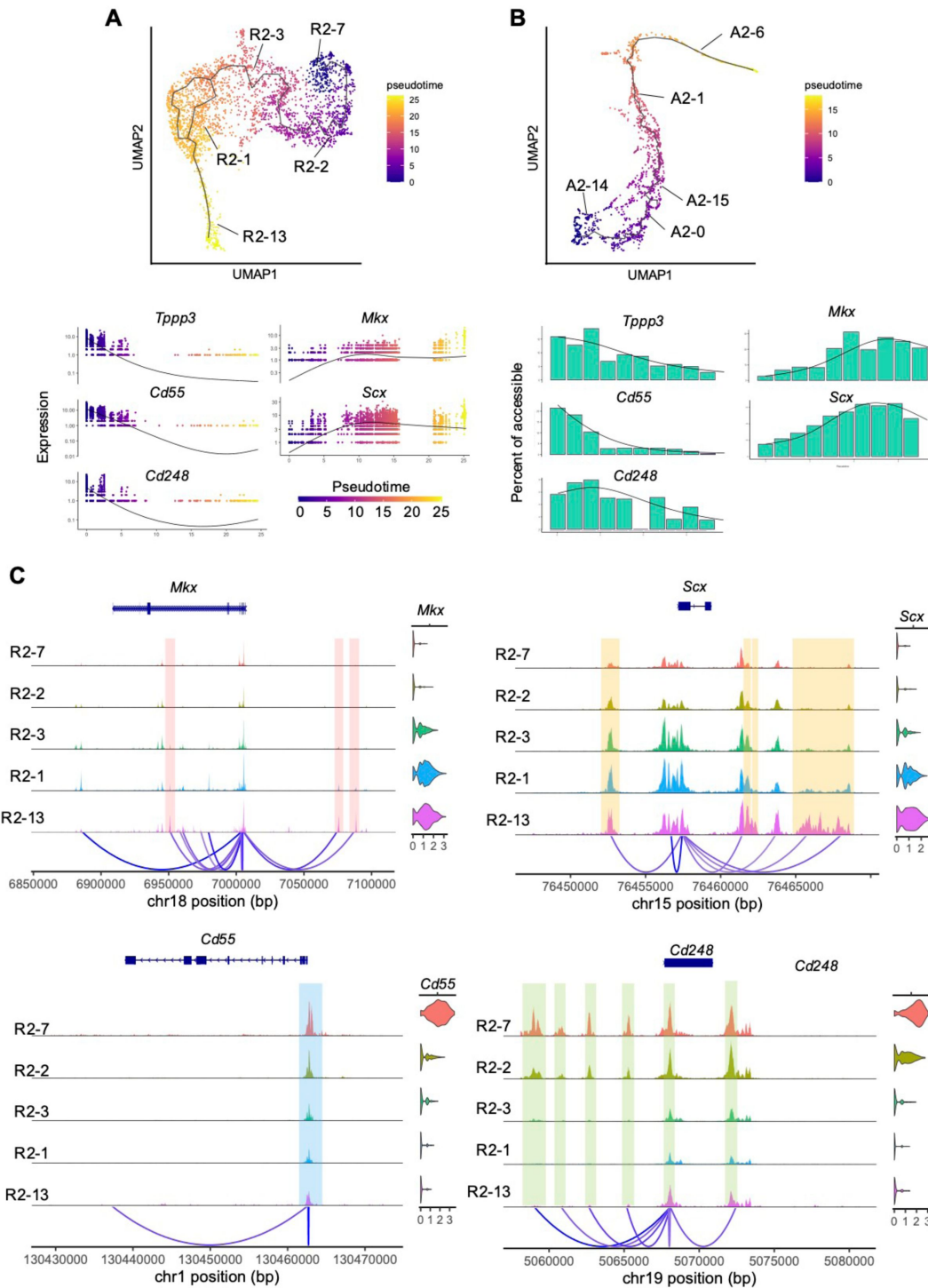


Figure 3.

Trajectory analysis and peak visualization of snRNA-seq and snATAC-seq data for the tendon and tendon stem/progenitor cell-related cluster.

(A) UMAP representation of snRNA-seq differentiation trajectory of tenocytes and TSPC lineage and pseudotime-dependent gene expression changes of *Tppp3*, *Cd55*, *Cd248*, *Mlx*, and *Scx*, as inferred using Monocle3.

(B) UMAP representation of snATAC-seq differentiation trajectory of tenocytes and the TSPC lineage and pseudotime-dependent gene expression changes, as inferred using Cicero.

(C) Coverage plots of *Mlx*, *Scx*, *Cd55*, and *Cd248*. Selected peaks that differ across each cluster are highlighted.

Comparison of Single Nucleus RNA+ATAC Analysis between 2-week and 6-week Mouse Achilles Tendon Cells

To confirm the presence of the identified *Cd55/Cd248* fraction at 6 weeks, we performed snRNA-seq and snATAC-seq on 6-week-old mouse Achilles tendon cells. Using the 2-week snATAC-seq data as a reference, we annotated the 6-week snATAC-seq clusters. Clusters A2-14 and A2-0 corresponded to clusters A6-12 and A6-0, respectively (**Supplemental Figure 4A, 4B** [↗](#)). Gene activity evaluation revealed that, similar to the 2-week data, the 6-week A6-12 and A6-0 clusters showed *Cd55/Cd248* high and *Cd34* dim, and *Cd55/Cd248* dim and *Cd34* high patterns, respectively. Furthermore, annotation of the 6-week snRNA-seq data using the 2-week snRNA-seq as a reference revealed clusters with expression patterns similar to those observed in snATAC-seq. In both cases, clusters with high *Cd55* and *Cd248* expression showed high gene activity (snATAC-seq) and gene expression (snRNA-seq) of *Tppp3*, *Pdgfra*, and *Ly6a*. These results confirm that the *Cd55* and *Cd248* high clusters identified in the 2-week snRNA-seq+snATAC-seq analysis are similarly detected at 6 weeks.

We also compared the snATAC-seq data between 2 and 6 weeks. No significant differences were observed in genomic annotations between the two time points (**Supplemental Figure 6A** [↗](#)). After merging the datasets (**Supplemental Figure 6B** [↗](#)), we compared gene activity of tendon/stem-related genes to evaluate changes in gene activity. We found that the activity of *Tppp3*, *Pdgfra*, *Ly6a*, *Cd55*, *Cd248*, and *Cd34* all decreased at 6 weeks (**Supplemental Fig 6C** [↗](#)). This was consistent with the decreased gene expression observed when comparing 2-week and 6-week data.

Estimation of Transcription Factor Activity in 2-week Mouse Achilles Tendon Cells

To estimate transcription factor activity in each cluster, we used the Single-Cell Regulatory Network Inference and Clustering (SCENIC) package (Lake et al., 2018) to calculate gene regulatory network activity from scRNA-seq gene expression data. SCENIC constructs gene expression networks centered on transcription factors and infers transcription factor activity in each cluster. We summarized the predicted transcription factor activities in tendon and stem/progenitor-related clusters R2-7, R2-2, R2-3, R2-1, and R2-13 identified by snRNA-seq (**Figure 4** [↗](#)).

Next, we evaluated the gene activity of these identified transcription factors in each snATAC-seq cluster, along with motif activity calculated by chromVAR (Schep et al., 2017). We also assessed the expression levels of transcription factors in each snRNA-seq cluster. In the most immature fractions, represented by cluster A2-14 in snATAC-seq and cluster R2-7 in snRNA-seq, KLF transcription factor 3 (*Klf3*), KLF transcription factor 4 (*Klf4*), and cAMP responsive element binding protein 5 (*Creb5*) showed consistent behavior in gene activity, motif activity, and gene expression. Additionally, Signal transducer and activator of transcription 2 (*Stat2*) and cAMP responsive element binding protein 3 like 1 (*Creb3l1*) showed high values in A2-0/A2-15 and A2-1, respectively. *Stat2* gene expression was not observed, likely due to false positives resulting from low expression levels. These results suggest a correlation between the transcription factor activity predicted by SCENIC and the gene activity and motif activity derived from snATAC-seq.

Furthermore, SCENIC can predict candidate transcription factors regulating each gene (**Table 1** [↗](#)). *Cd55* and *Cd248* were predicted to be regulated by *Klf3* and *Klf4*, while *Mkx* and *Scx* were predicted to be under the control of *Creb3l1*. These predictions were consistent with the data calculated from gene activity and motif activity in snATAC-seq.

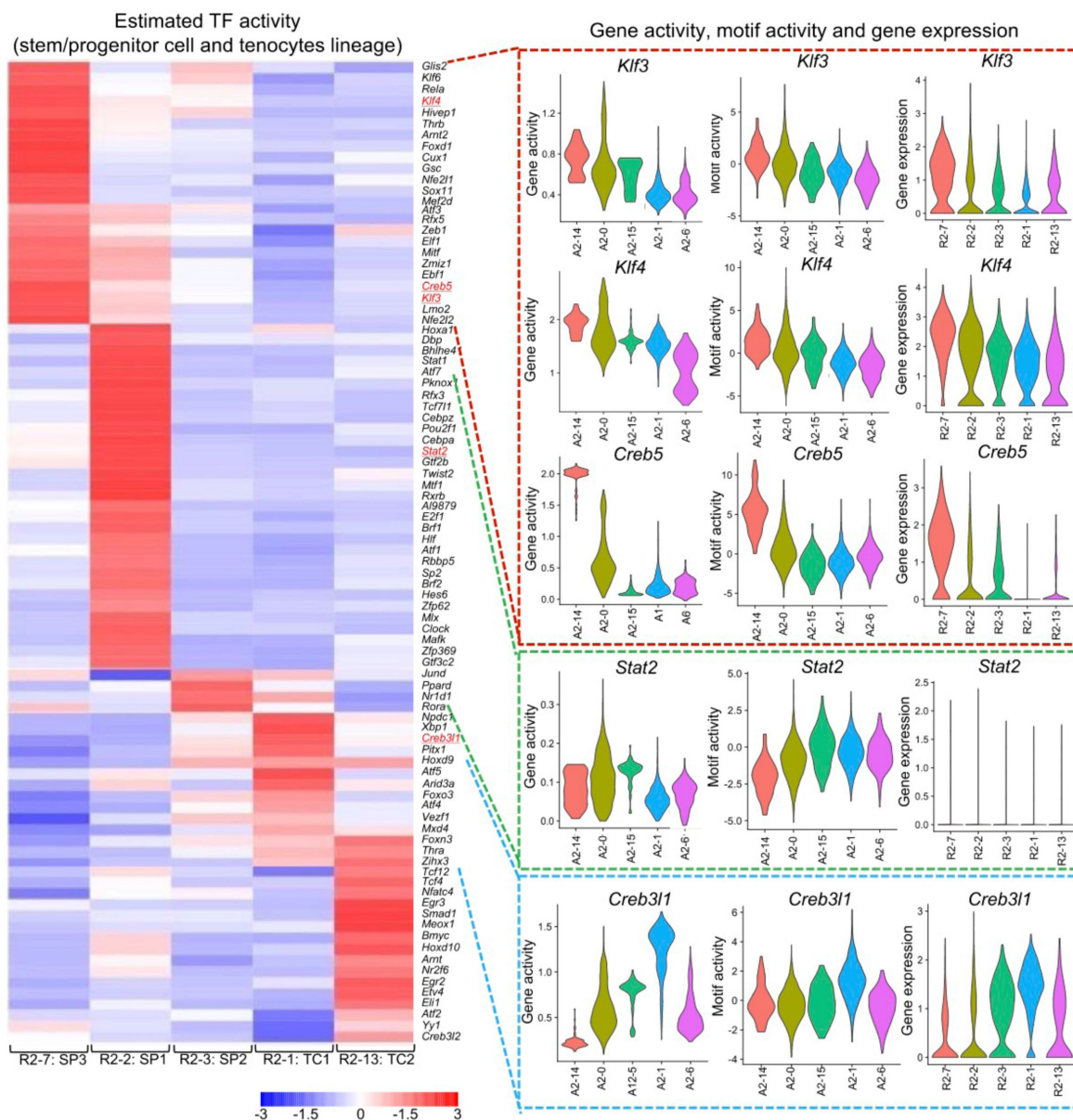


Figure 4.

Transcription factor landscapes of 2-week mouse Achilles tendons.

SCENIC analysis of transcription factor activity based on 2-week snRNA-seq data for tenocytes and the TSPC lineage (Left). Validation was performed based on the gene activity and motif activity of 2-week snATAC-seq and gene expression of 2-week snRNA-seq (right).

Estimated TF activity : Cd55		Estimated TF activity : Cd248		Estimated TF activity : Mlx		Estimated TF activity : Scx	
TF	spearman correlation	TF	spearman correlation	TF	spearman correlation	TF	spearman correlation
Klf4	0.264195541	Hic1	0.375565336	Creb3l1	0.320638653	Creb3l1	0.316522613
Klf3	0.262413411	Klf3	0.277768834	Zfhx3	0.169341424	Npdc1	0.163452737
Zeb1	0.219593135	Zmiz1	0.272991765	Mxi1	0.149048856	Ets2	0.145463784
Pbx1	0.204085218	Klf4	0.260526766	Bhlhe40	0.144301452	Mix1	0.121248221
Hic1	0.202481054	Zeb1	0.250583629	Ets2	0.137445497	Bhlhe41	0.115930755
Klf2	0.196232716	Sp3	0.2502976	Max	0.116134957	Elk3	0.10803805
Nfe2l2	0.196076187	Kdm5b	0.193229132	Elk3	0.098325126	Bmyc	0.104968457

Table 1

Estimated TF activity in each gene

In vitro Evaluation of CD55 and CD248

Immunostaining evaluation of CD55 and CD248, the identified candidate stem/progenitor markers, revealed expression in the tendon sheath (**Figure 5C** [↗](#), 5A). This was consistent with previous reports of localized expression of *Tppp3/Pdgfra*-positive cells in the tendon sheath and our analysis showing co-expression of *Tppp3/Pdgfra* and *Cd55/Cd248*. Given that *Cd55* and *Cd248* expression appeared to reflect *Tppp3*, *Pdgfra*, and *Ly6a* expression more sensitively than *Cd34*, we extracted tendon cells from 2-week-old mice and sorted them by FACS to determine the biological phenotype of *Cd55* and *Cd248* positive cells (**Figure 5B** [↗](#)).

Gene expression analysis of sorted cells showed that CD55/CD248 positive cells, compared to negative cells, had lower expression of *Mkx*, *Scx*, *Col1a1*, and *Creb3l1*, but higher expression of *Ly6a*, *Tppp3*, *Pdgfra*, *Creb5*, and *Klf3*, consistent with snRNA-seq analysis (**Supplemental Figure 6** [↗](#)). To assess stem/progenitor capacity, we performed colony formation assays. CD55/CD248 sorted cells showed significantly increased colony formation compared to CD55/CD248 negative cells (control) (**Figure 5C** [↗](#)).

We then evaluated the differentiation tendencies of these cells towards tendon, cartilage, and bone. Tendon differentiation resulted in clusters of spindle-shaped cells from CD55/CD248 positive cells, suggesting tenogenic differentiation (**Figure 5D** [↗](#)). Gene expression analysis revealed that while CD55/CD248 positive cells initially had lower *Mkx* and *Scx* expression than negative cells, this pattern reversed after differentiation (**Figure 5E** [↗](#)). Increased expression of other tendon-related genes such as *Col1a1* and *Bgn* was also observed. In contrast, no significant increases in *Sox9* and Collagen type II alpha 1 chain (*Col2a1*)(chondrogenic) or RUNX family transcription factor 2 (*Runx2*), and Alkaline phosphatase, biomineralization associated (*Alpl*) (osteogenic) expression were observed compared to negative cells during cartilage and bone differentiation assays (**Supplemental Figure 7** [↗](#)). These results suggest that CD55/CD248 positive cells possess tenogenic differentiation capacity.

To further evaluate the tenogenic potential of CD55/CD248 positive cells, we created tendon-like tissue (bio-tendon) using our previously reported 3D stretch stimulation culture system. SEM and TEM analysis to assess collagen fiber density and thickness showed that bio-tendons derived from CD55/CD248 positive cells had an increased proportion of collagen fibers parallel to the stretch direction and increased collagen fiber diameter. TEM also revealed characteristic banding patterns and triple helix structures in these bio-tendons, indicating mature collagen organization (Wieczorek et al., 2015) (**Figure 5F-H** [↗](#)). Stretch tests to evaluate the mechanical capacity of the bio-tendons showed high tensile strength (**Figure 5I** [↗](#)).

These results demonstrate that CD55/CD248 positive cells have a tendency to differentiate into tendon cells.

Discussion

Tendons are known for their low cellular content, making complete functional recovery after injury challenging and increasing the risk of re-rupture. As a result, cell therapy has garnered attention as a novel treatment strategy, distinct from current conservative and surgical approaches. However, this requires a thorough understanding of Tendon Stem/Progenitor Cells (TSPCs). Research on TSPCs has been limited due to the lack of identified characteristic surface antigens.

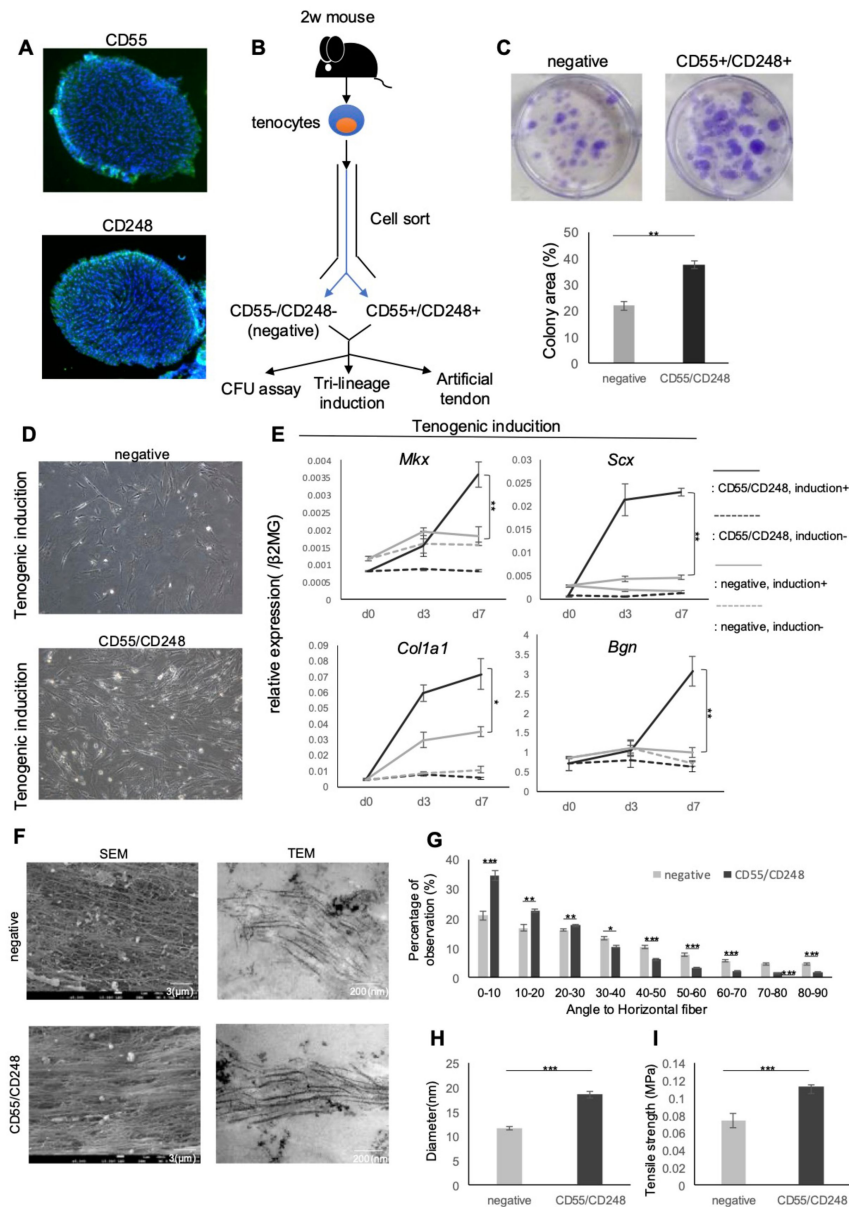


Figure 5.

***In vitro* analysis of CD55+/CD248+ TSPCs.**

- (A) Immunohistochemical image of 2-week mouse Achilles tendons. CD55 and CD248, green; DAPI, blue.
- (B) Schema of the *in vitro* assessment of the capacity of CD55+/CD248+ TSPCs as the differentiation toward tenocytes.
- (C) Colony-forming efficiency of CD55+/CD248+ and CD55-/CD248- (negative) TSPCs. Colonies were stained with crystal violet ($n = 6$). CD55+/CD248+ TSPC exhibited higher clonogenic capacity. Data are presented as means $\pm$ SEM. $^{**}p < 0.01$.
- (D) Morphological changes of CD55+/CD248+ and negative TSPCs after tenogenic induction.
- (E) Quantitative PCR of tendon-related gene expression in CD55+/CD248+ and negative TSPCs after tenogenic induction ($n = 3$). Data are presented as means $\pm$ SD. $^{**}p < 0.01$, $^{*}p < 0.05$.
- (F) SEM and TEM imaging of artificial tendons derived from CD55+/CD248+ and negative TSPCs. Data are presented as means $\pm$ SD. $^{**}p < 0.01$, $^{*}p < 0.05$.
- (G) Proportions of fiber alignment for each artificial tendon ($n = 4$). Data are presented as means $\pm$ SEM. $^{***}p < 0.005$, $^{**}p < 0.01$, $^{*}p < 0.05$.
- (H) Diameter of collagen fiber in each artificial tendon based on TEM imaging ($n = 4$). Data are presented as means $\pm$ SD. $^{***}p < 0.005$.
- (I) Tensile strength (MPa) of each artificial tendon ($n = 5$). Data are presented as means $\pm$ SEM. $^{***}p < 0.005$.

To our knowledge, no multi-omics analysis comparing juvenile and mature mouse tendon cells has been conducted. In this study, we performed single-cell RNA-seq along with snRNA-seq+snATAC-seq on nuclei isolated from 2-week and 6-week mouse Achilles tendons. This approach allowed us to identify candidate surface antigens for TSPCs and elucidate the dynamism of transcription factors involved in post-developmental tendon growth. As a result, we discovered a novel combination of *Cd55* and *Cd248* as surface antigens showing characteristic expression in the TSPC fraction. While scRNA-seq analysis also showed *Cd34* expression characteristic of the TSPC fraction, consistent with previous reports, snRNA-seq + snATAC-seq analysis revealed that the *Cd34* high-expression cluster could be further classified into two clusters based on *Cd55* and *Cd248* expression patterns. The cluster showing high expression of both *Cd55* and *Cd248* also exhibited high expression of *Tppp3*, *Pdgfra*, and *Ly6a*, suggesting that isolating *Cd55* and *Cd248* positive fractions may more sensitively capture immature populations compared to *Cd34*.

snRNA-seq+snATAC-seq analysis provided simultaneous information on gene expression levels and open chromatin regions for each cluster. Notably, *Cd248* showed several peaks upstream of the TSS in high-expression clusters, likely reflecting its expression regulation mechanism. We also identified several peaks that increased proportionally with expression for *Mkx* and *Scx*. Given the many unknowns in *Mkx* and *Scx* expression regulation mechanisms (Guerquin et al., 2013; Otabe et al., 2015), we used the SCENIC package to predict upstream transcription factors based on expressed genes in each snRNA-seq cluster. Combined with gene activity and motif activity data from snATAC-seq, *Creb3l1* was identified as a transcription factor regulating *Mkx* and *Scx* expression. *Creb3l1* has been reported to increase in expression throughout development (Liu et al., 2015 [DOI](#)). *Klf3*, *Klf4*, and *Creb5* were identified as characteristic transcription factors in the TSPC fraction. Recent reports have highlighted the role of the *Klf* family in limb development (Kult et al., 2021), suggesting its potential importance in tendon differentiation. *Creb5* has been reported to show increased expression from E11 to E13 (Liu et al., 2015 [DOI](#)), suggesting a role in early developmental stages. Furthermore, *Creb5* has been reported to regulate Proteoglycan 4 (*Prg4*) expression (Zhang et al., 2021). Further investigation into the functions of these transcription factors in TSPCs is necessary.

CD55 was identified by Hoffmann et al. in 1969 as a surface antigen functioning as a complement inhibitory factor on erythrocytes. CD55 is known to be expressed from early developmental stages and characterizes the initial differentiation stage of hematopoietic stem cells (Guo et al., 2013). It is also expressed in MSCs and has been reported as an early progenitor marker for mouse mammary epithelial cells (Pal et al., 2017). CD248 was identified in 1992 as an antigen for the FB5 antibody reacting with vascular wall cells (Rettig et al., 1992). CD248 is a transmembrane glycoprotein expressed in pericytes and fibroblasts during developmental stages. Regarding their relationship, CD55 and CD248 have been reported to be expressed in stromal cells during the early stages of arthritis (Choi et al., 2017), but their detailed mechanisms in tendons remain unclear.

Previously suggested TSPC surface antigen candidates such as *Cd73*, *Cd90*, and *Cd105* showed poor correlation with the expression of genes like *Tppp3*. Moreover, the initial report on TSPCs (Bi et al., 2007 [DOI](#)) described TSPCs as *Scx*+*Cd34*-. The high expression of CD55 and CD248 in the tendon sheath, similar to *Tppp3* (Harvey et al., 2019 [DOI](#); Staverosky et al., 2019 [DOI](#)), suggests the possibility of cells with tenogenic differentiation potential exhibiting *Scx*+*Cd34*-expression patterns within the tendon. Future studies using lineage tracing experiments with mice labeled for CD55 and CD248 are necessary to analyze the developmental functions of CD55 and CD248 positive cells and their roles in injury healing in more detail.

Clinically, to our knowledge, few studies have sorted TSPCs based on surface antigens and examined their *in vitro* tendon tissue-generating ability. In this study, we found that cells sorted for CD55/CD248 showed higher clonogenicity compared to CD55-/CD248-cells and demonstrated

superior tendon tissue-generating ability in an artificial tendon model. In the future, CD55/CD248 double-positive TSPC cells may prove useful in clinical applications such as *in vitro* artificial tendon creation (Tsutsumi et al., 2022 [DOI](#)) and cell therapy for tendon injuries (Huang et al., 2021).

Data availability

FASTQ data of RNA-Seq and ATAC-seq are deposited in DDBJ under accession number PRJDB18857.

Acknowledgements

We thank all the members of the Department of Systems BioMedicine at Institute of Science Tokyo for their support. We also thank the Research Core at Institute of Science Tokyo for supporting cell sorting.

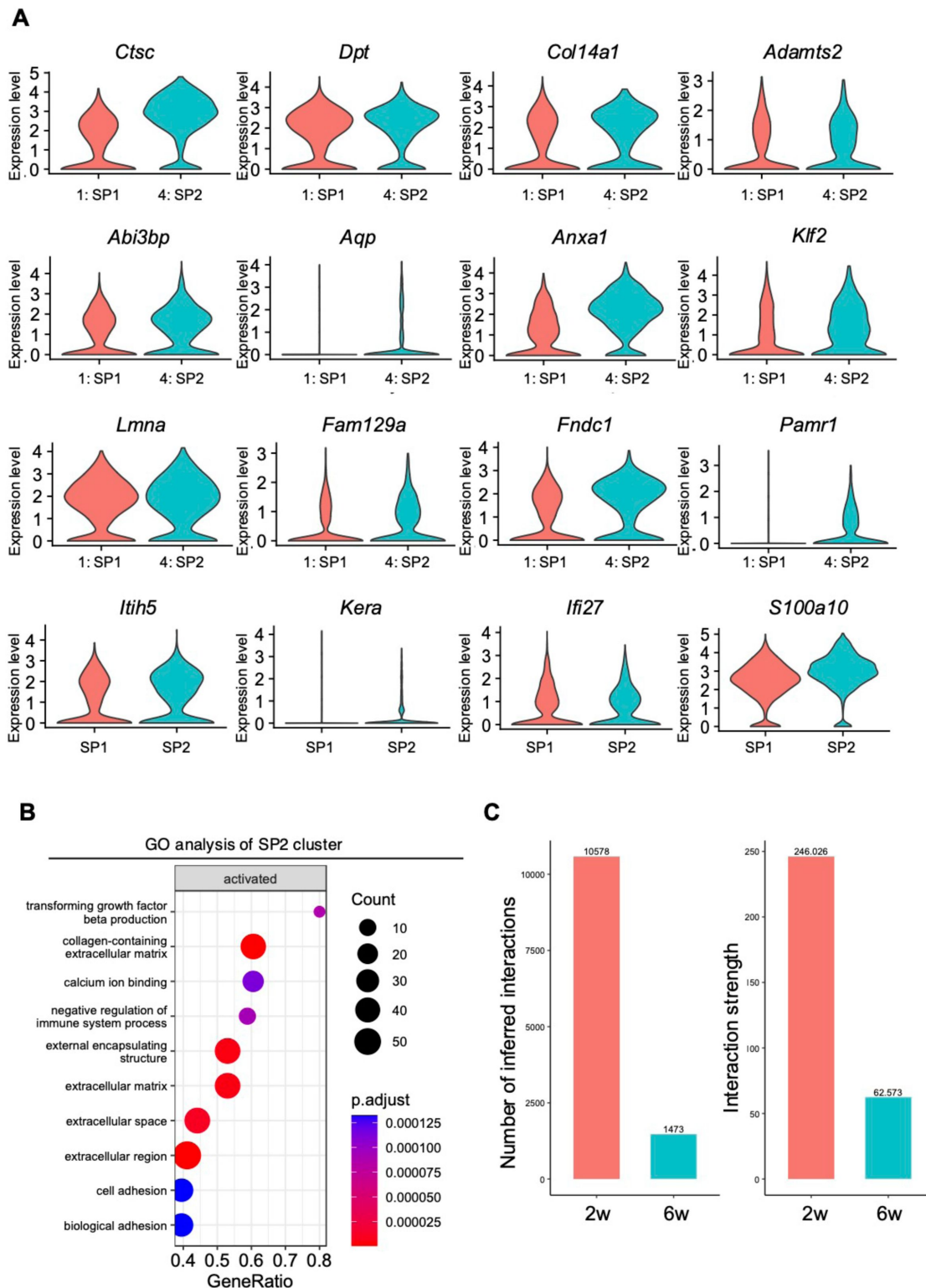
Additional information

Author contributions

Conceptualization, H.T., T.C., and H.A.; Methodology, H.T., T.C., and H.A.; Formal analysis, H.T.; Investigation, H.T., F. Y., T. M., T. K., A. K., K. A., Y. S.; Writing – Original Draft, H.T.; Writing – Review & Editing, T.C., and H.A.; Funding Acquisition, H.A.; Supervision, H.A.

Funding

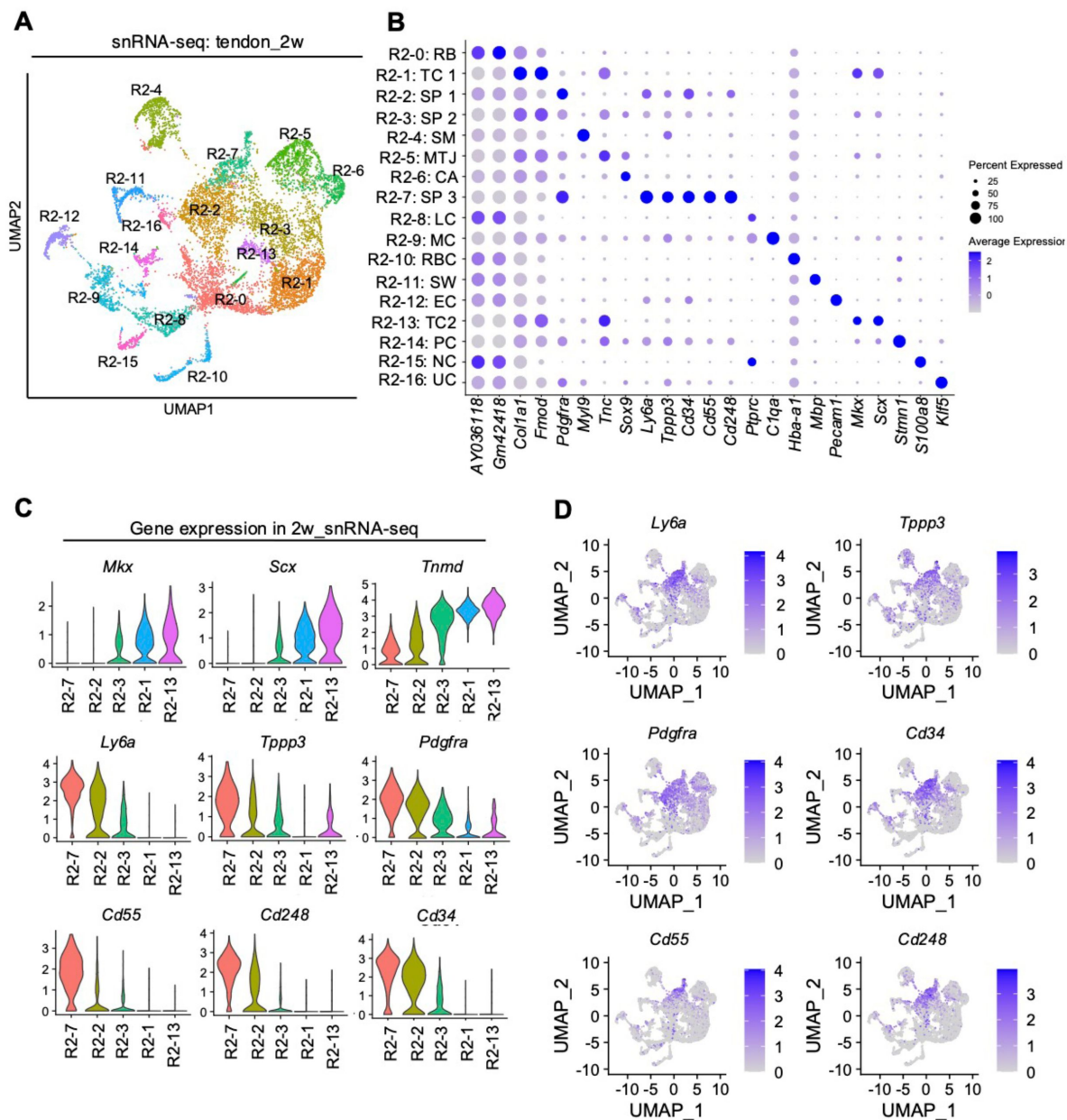
This work was supported by JSPS KAKENHI (Grant Numbers JP15H02560, JP20H05696, 16H06279 (PAGS)), AMED (Grant Numbers JP21gm0810008, JP23ym0126805, JP24gm0010009), and NIH (Grant Number R01AR080127) to H.A.



Supplemental Figure 1.

Comparison of 2w- and 6w-scRNA seq results.

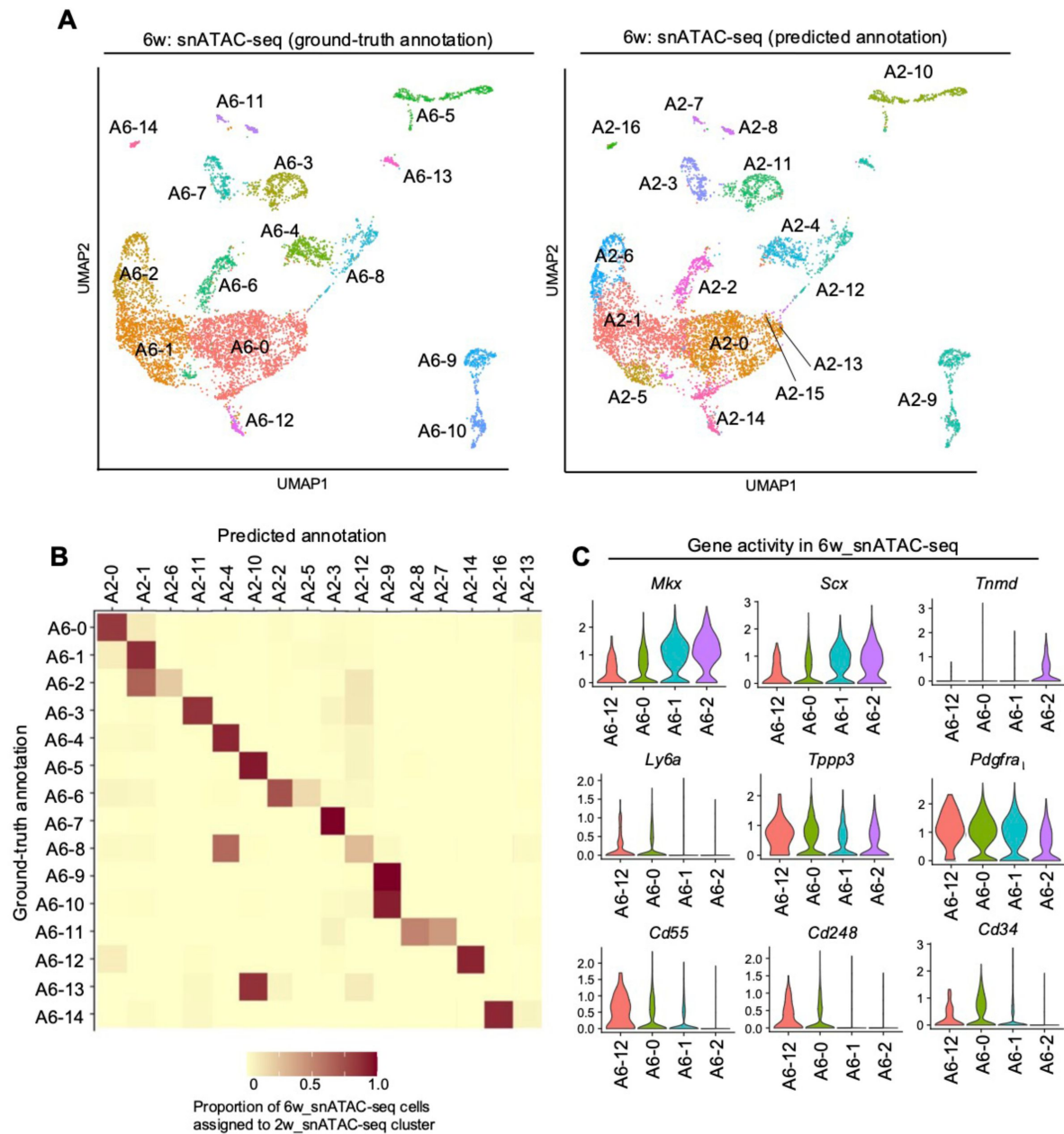
- (A) Violin plot of gene expression enriched in the limb bud (Development (2014) 141, 3683-3696) in each cluster.
 (B) Gene Ontology (GO) terms associated with genes with upregulated expression in the SP2 cluster.
 (C) Number of inferred interactions and interaction strength of genes expressed within 2w- and 6w-mouse scRNA-seq data.



Supplemental Figure 2.

Analysis of 2-week snRNA-seq data.

- (A) UMAP snRNA-seq clustering of cells harvested from 2-week mouse Achilles tendons.
- (B) Dot plot of average gene expression levels of the indicated genes in 2-week snRNA-seq clusters. RB, ribosomal RNA; TC1, tenocyte_1; SP1, stem/progenitor cell_1; SP2, stem/progenitor cell_2; SM, smooth muscle cell; MTJ, myotendinous junction cell; CA, cartilage; SP3, stem/progenitor cell_3; LC, lymphocyte; MC, macrophage; RBC, red blood cell; SW, Schwann cell; EC, endothelial cell; TC2, tenocyte_2; PC, proliferating cell; NC, neutrophil; UC, unspecified cell.
- (C) Violin plot of tenocytes and TSPC-related gene expression in each cluster.
- (D) Feature plot of TSPC-related gene expression.



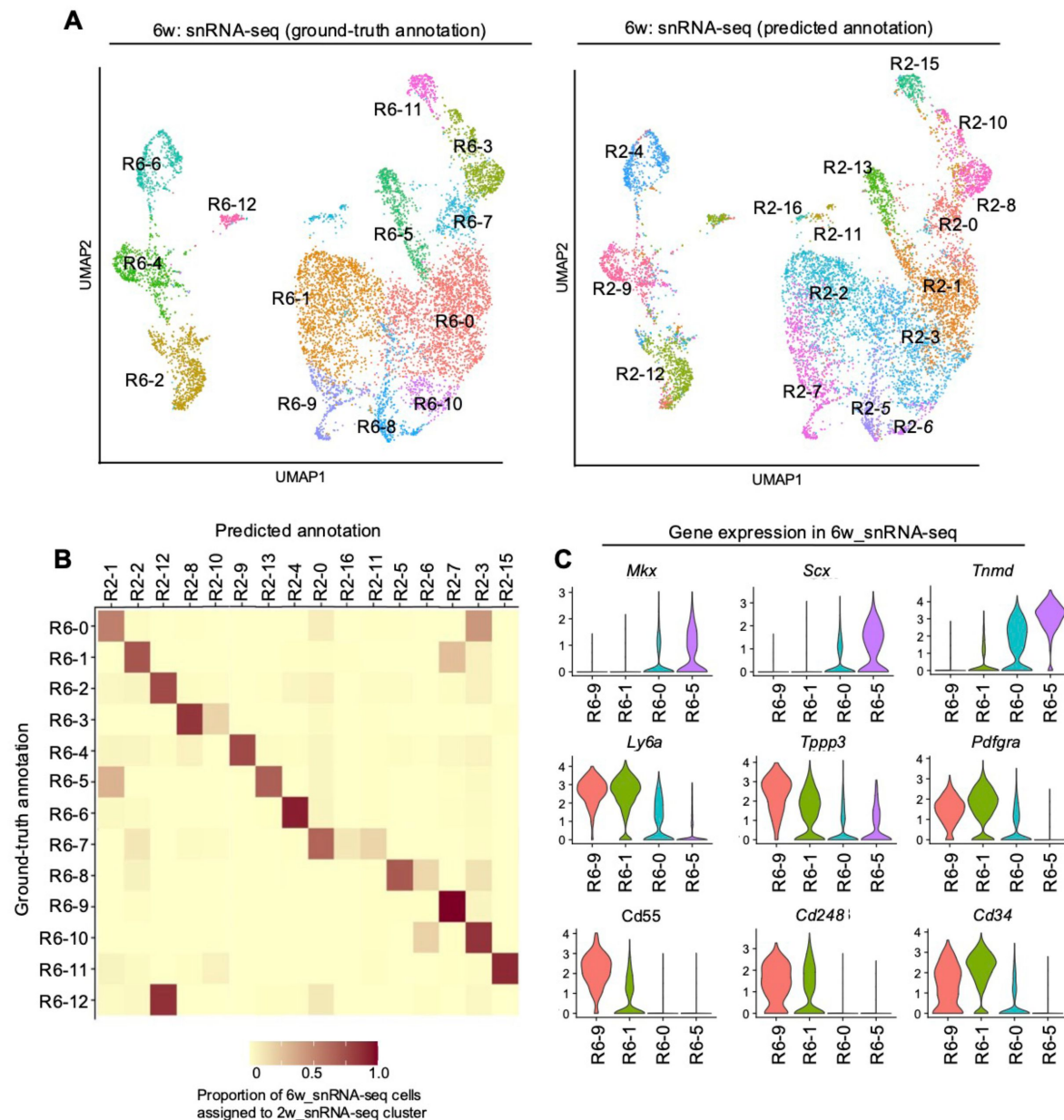
Supplemental Figure 3.

Analysis of 6-week snATAC-seq.

(A) UMAP snATAC-seq clustering of cells harvested from 6-week mouse Achilles tendons. Annotation was based on gene activity (ground-truth annotation, left) and the predicted annotation inferred from 2-week snATAC-seq (right).

(B) Identification of matching cell clusters between the 2-week and 6-week snATAC-seq data, visualized as a heatmap.

(C) Violin plot of tenocytes and TSPC-related gene activity in each cluster.



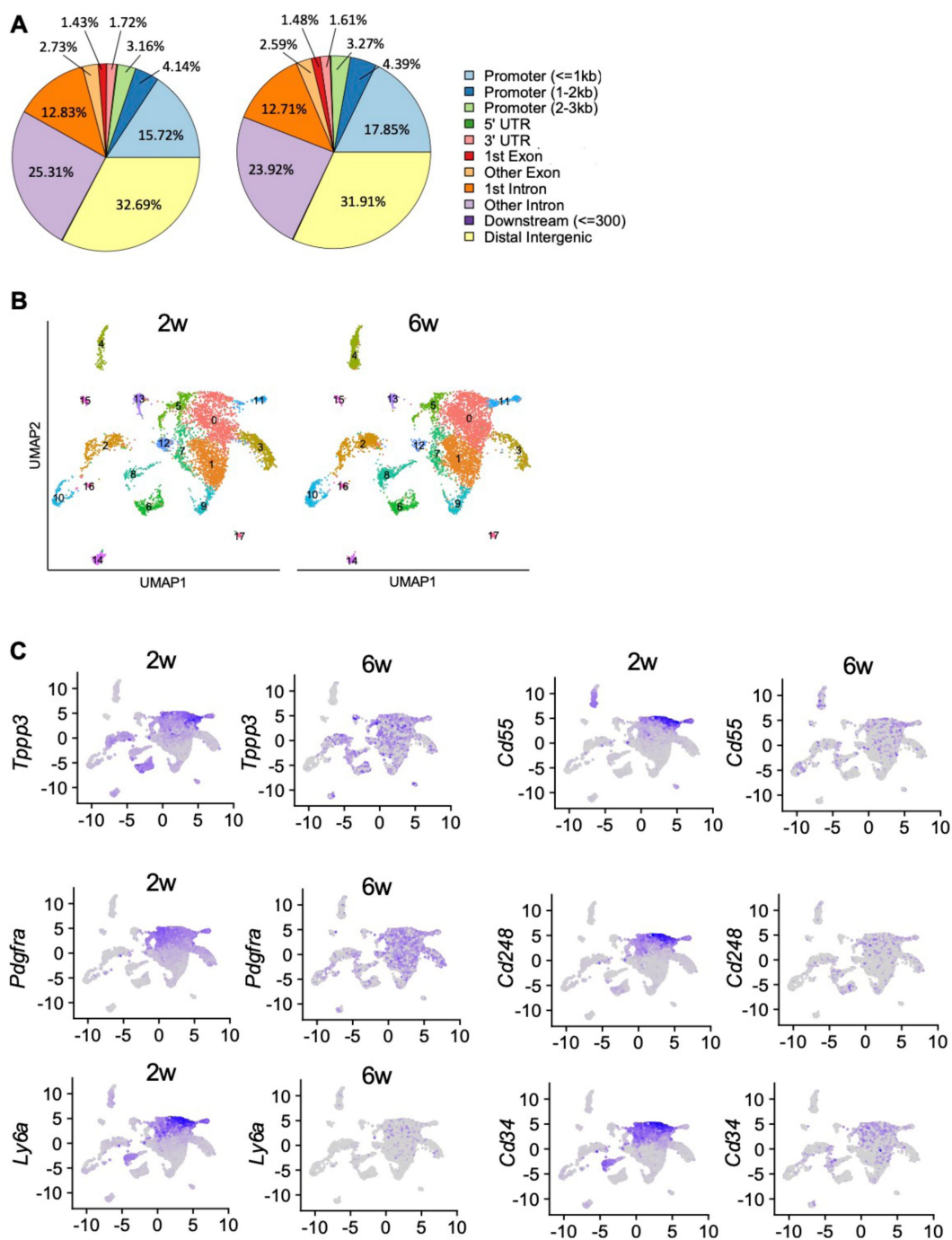
Supplemental Figure 4.

Analysis of 6-week snRNA-seq.

(A) UMAP snRNA-seq clustering of cells harvested from 6-week mouse Achilles tendons. Annotation was based on each gene expression (ground-truth annotation, left) and predicted annotation inferred from 2-week snRNA-seq (right).

(B) Identification of matching cell clusters between the 2-week and 6-week snRNA-seq data visualized as a heatmap.

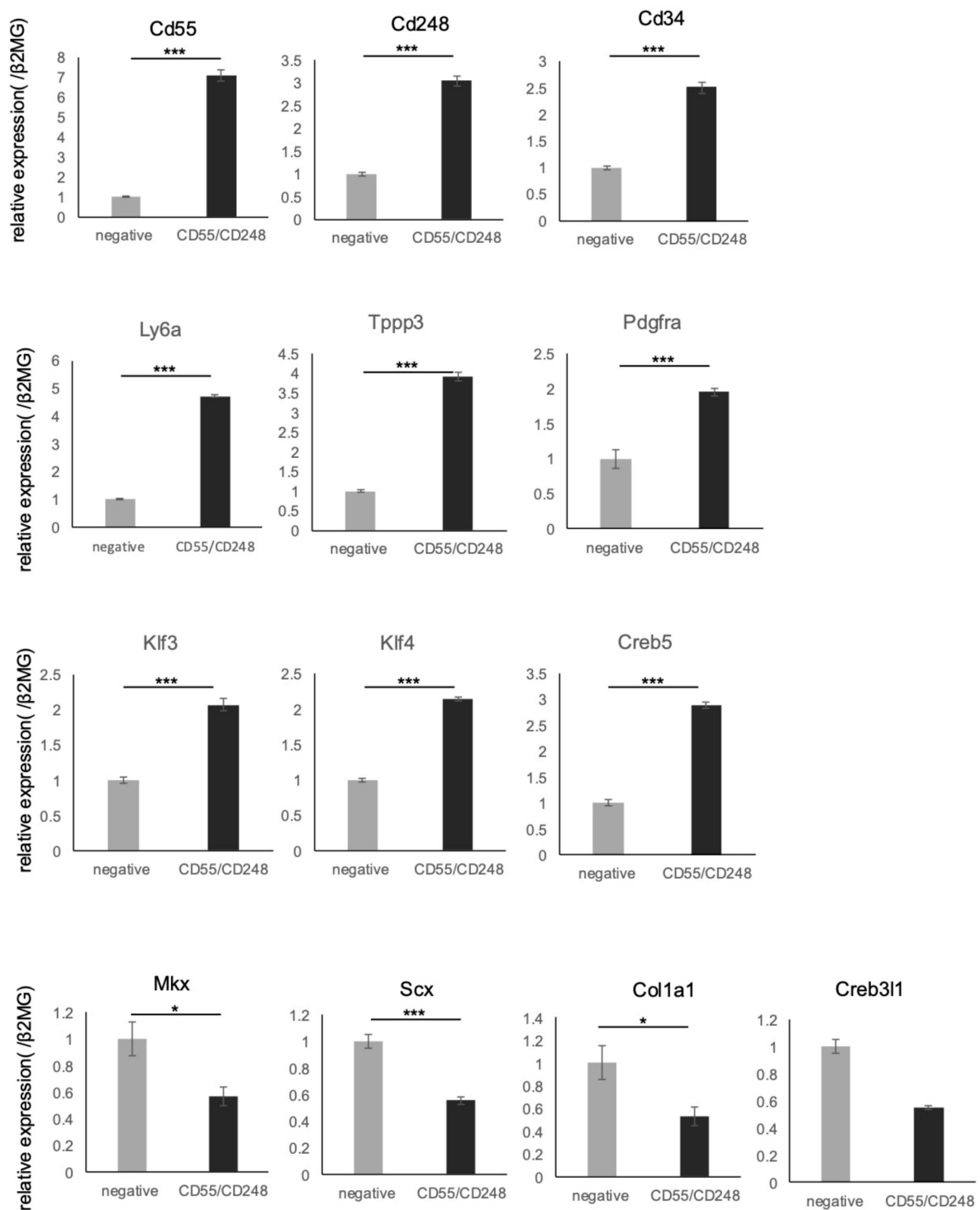
(C) Violin plot of tenocytes and TSPC-related gene expression in each cluster.



Supplemental Figure 5.

Comparison of 2-week and 6-week snATAC-seq.

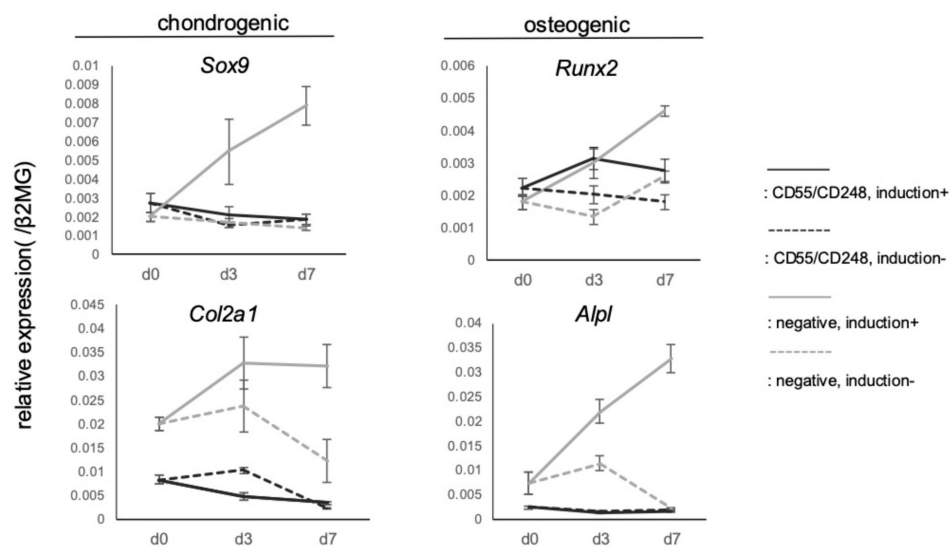
- (A) Circle plot of annotated differentially accessible regions for each data.
 (B) Integrated UMAP snATAC-seq clustering of cells harvested from 2-week and 6-week mouse Achilles tendons.
 (C) Feature plot of TSPC-related gene activity in each dataset.



Supplemental Figure 6.

Gene expression changes in CD55+/CD248+ and negative TSPCs.

Quantitative PCR of gene expression in CD55+/CD248+ and negative TSPCs (n = 4). Data are presented as means $\pm$ SEM. * p < 0.05, *** p < 0.005.



Supplemental Figure 7.

Chondrogenic and osteogenic induction of CD55+/CD248+ and negative TSPCs.

Quantitative PCR of tendon-related genes in CD55+/CD248+ and negative TSPCs after chondrogenic and osteogenic induction (n = 4). Data are presented as means $\pm$ SEM. ** $p < 0.01$, * $p < 0.05$.

References

- Andarawis-Puri N., Flatow EL., Soslowsky LJ 2015) **Tendon basic science: Development, repair, regeneration, and healing** *J Orthop Res* **33**:780–4
- Asahara H., Inui M., Lotz MK 2017) **Tendons and Ligaments: Connecting Developmental Biology to Musculoskeletal Disease Pathogenesis** *J Bone Miner Res* **32**:1773–1782
- Becht E., McInnes L., Healy J., Dutertre CA., Kwok IWH., Ng LG., Ginhoux F., Newell EW. 2019) **Dimensionality reduction for visualizing single-cell data using UMAP** *Nat Biotechnol* **37**:38–44
- Bi Y., Ehrichtiou D., Kilts TM, Inkson CA, Embree MC., Sonoyama W., Li L., Leet AI., Seo BM., Zhang L., Shi S., Young MF 2007) **Identification of tendon stem/progenitor cells and the role of the extracellular matrix in their niche** *Nat Med* **13**:1219–27
- Butler A., Hoffman P., Smibert P., Papalexi E., Satija R 2018) **Integrating single-cell transcriptomic data across different conditions, technologies, and species** *Nat Biotechnol* **36**:411–420
- Cao Y., Liu Y., Liu W., Shan Q., Buonocore SD., Cui L 2002) **Bridging tendon defects using autologous tenocyte engineered tendon in a hen model** *Plast Reconstr Surg* **110**:1280–9
- Chen B., Wang B., Zhang WJ., Zhou G., Cao Y., Liu W 2012) **In vivo tendon engineering with skeletal muscle derived cells in a mouse model** *Biomaterials* **33**:6086–97
- Chen L., Liu JP., Tang KL., Wang Q., Wang GD., Cai XH., Liu XM 2014) **Tendon derived stem cells promote platelet-rich plasma healing in collagenase-induced rat achilles tendinopathy** *Cell Physiol Biochem* **34**:2153–68
- Cho YT., Hsu WY., Lin LF., Lin YN 2018) **Kinesio taping reduces elbow pain during resisted wrist extension in patients with chronic lateral epicondylitis: a randomized, double-blinded, cross-over study** *BMC Musculoskelet Disord* **19**:193
- Connizzo BK., Yannascoli SM., Soslowsky LJ 2013) **Structure-function relationships of postnatal tendon development: a parallel to healing** *Matrix Biol* **32**:106–16
- Craft A. M., Ahmed N., Rockel J. S., Baht G. S., Alman B. A., Kandel R. A., Grigoriadis A. E., Keller G. M 2013) **Specification of chondrocytes and cartilage tissues from embryonic stem cells** *Development* **140**:2597–2610
- Diez-Roux G., Banfi S., Sultan M., Geffers L., Anand S., Rozado D., Magen A., Canidio E., Pagani M., Peluso I., Lin-Marq N., Koch M., Bilio M., Cantiello I., Verde R., De Masi C., Bianchi SA., Cicchini J., Perroud E., Mehmeti S., Dagand E., Schrinner S., Nürnberger A., Schmidt K., Metz K., Zwingmann C., Brieske N., Springer C., Hernandez AM., Herzog S., Grabbe F., Sieverding C., Fischer B., Schrader K., Brockmeyer M., Dettmer S., Helbig C., Alunni V., Battaini MA., Mura C., Henrichsen CN., Garcia-Lopez R., Echevarria D., Puellas E., Garcia-Calero E., Kruse S., Uhr M., Kauck C., Feng G., Milyaev N., Ong CK., Kumar L., Lam M., Semple CA., Gyenesi A., Mundlos S., Radelof U., Lehrach H., Sarmientos P., Reymond A., Davidson DR., Dollé P., Antonarakis SE.,

- Yaspo ML., Martinez S., Baldock RA., Eichele G., Ballabio A. 2011) **A High-Resolution Anatomical Atlas of the Transcriptome in the Mouse Embryo** *PLoS Biol* **9**:e1000582
- Gracey E., Burssens A., Cambré I., Schett G., Lories R., McInnes IB., Asahara H., Elewaut D 2020) **Tendon and ligament mechanical loading in the pathogenesis of inflammatory arthritis** *Nat Rev Rheumatol* **16**:193–207
- Fang F., Xiao Y., Zelzer E., Leong KW., Thomopoulos S 2022) **A mineralizing pool of Gli1-expressing progenitors builds the tendon enthesis and demonstrates therapeutic potential** *Cell Stem Cell* **29**:1669–1684
- Franken NA., Rodermond HM., Stap J., Haveman J., van Bree C. 2006) **Clonogenic assay of cells in vitro** *Nat Protoc.* **2006** **1**:2315–9
- Grinstein M., Dingwall HL., O'Connor LD., Zou K., Capellini TD., Galloway JL 2019) **A distinct transition from cell growth to physiological homeostasis in the tendon** *eLife* **8**:e48689
- Harvey T., Flamenco S., Fan CM 2019) **A Tppp3 + Pdgfra + tendon stem cell population contributes to regeneration and reveals a shared role for PDGF signalling in regeneration and fibrosis** *Nat Cell Biol* **21**:1490–1503
- Guilak F., Butler DL., Goldstein SA., Baaijens FP 2014) **Biomechanics and mechanobiology in functional tissue engineering** *J Biomech* **47**:1933–40
- Gumucio JP., Sugg KB., Mendias CL 2015) **TGF- β superfamily signaling in muscle and tendon adaptation to resistance exercise** *Exerc Sport Sci Rev* **43**:93–9
- Guzmán C., Bagga M., Kaur A., Westermarck J., Abankwa D 2014) **ColonyArea: an ImageJ plugin to automatically quantify colony formation in clonogenic assays** *PLoS One* **9**:e92444
- Havis E., Bonnin MA., Olivera-Martinez I., Nazaret N., Ruggiu M., Weibel J., Durand C., Guerquin MJ., Bonod-Bidaud C., Ruggiero F., Schweitzer R., Duprez D 2014) **Transcriptomic analysis of mouse limb tendon cells during development** *Development* **141**:3683–96
- Hittinger M., Czyz ZT., Huesemann Y., Maneck M., Botteron C., Kaeufl S., Klein CA., Polzer B 2013) **Molecular profiling of single Sca-1+/CD34+,-cells--the putative murine lung stem cells** *PLoS One* **8**:e83917
- Hoffmann EM 1969) **Inhibition of complement by a substance isolated from human erythrocytes. II. Studies on the site and mechanism of action** *Immunochemistry* **6**:405–19
- Holmes C., Stanford WL 2007) **Concise review: stem cell antigen-1: expression, function, and enigma** *Stem Cells* **25**:1339–47
- Howell K., Chien C., Bell R., Laudier D., Tufa SF., Keene DR., Andarawis-Puri N., Huang AH 2017) **Novel Model of Tendon Regeneration Reveals Distinct Cell Mechanisms Underlying Regenerative and Fibrotic Tendon Healing** *Sci Rep* **7**:45238
- Isola JVV., Ocañas SR., Hubbard CR., Ko S., Mondal SA., Hense JD., Carter HNC., Schneider A., Kovats S., Alberola-Ila J., Freeman WM., Stout MB 2024) **A single-cell atlas of the aging mouse ovary** *Nat Aging* **4**:145–162
- Jozsa L., Kannus P., Balint JB., Reffy A 1991) **Three-dimensional ultrastructure of human tendons** *Acta Anat (Basel)* **142**:306–12

- Kendal AR., Layton T., Al-Mossawi H., Appleton L., Dakin S., Brown R., Loizou C., Rogers M., Sharp R., Carr A 2020) **Multi-omic single cell analysis resolves novel stromal cell populations in healthy and diseased human tendon** *Sci Rep* **10**:13939
- Leong DJ., Sun HB 2016) **Mesenchymal stem cells in tendon repair and regeneration: basic understanding and translational challenges** *Ann. N. Y. Acad. Sci* **1383**:88–96
- Leong NL., Kator JL., Clemens TL., James A., Enamoto-Iwamoto M., Jiang J 2020) **Tendon and Ligament Healing and Current Approaches to Tendon and Ligament Regeneration** *J Orthop Res* **38**:7–12
- Li Y., Wu T., Liu S 2021) **Identification and Distinction of Tenocytes and Tendon-Derived Stem Cells** *Front Cell Dev Biol* **9**:629515
- Liu H., Xu J., Liu CF., Lan Y., Wylie C., Jiang R 2015) **Whole transcriptome expression profiling of mouse limb tendon development by using RNA-seq** *J Orthop Res* **33**:840–848
- Loiacono C., Palermi S., Massa B., Belviso I., Romano V., Gregorio AD., Sirico F., Sacco AM 2019) **Tendinopathy: Pathophysiology, Therapeutic Options, and Role of Nutraceuticals. A Narrative Literature Review** *Medicina (Kaunas)* **55**:447
- Lui PP 2015) **Markers for the identification of tendon-derived stem cells in vitro and tendon stem cells in situ - update and future development** *Stem Cell Res Ther* **6**:106
- Lui PPY., Wong CM 2019) **Biology of Tendon Stem Cells and Tendon in Aging** *Front Genet* **10**:1338
- McGinnis CS., Murrow LM., Gartner ZJ 2019) **DoubletFinder: Doublet Detection in Single-Cell RNA Sequencing Data Using Artificial Nearest Neighbors** *Cell Syst* **8**:329–337
- Mendias CL., Gumucio JP., Bakhurin KI., Lynch EB., Brooks SV 2012) **Physiological loading of tendons induces scleraxis expression in epitenon fibroblasts** *J Orthop Res* **30**:606–12
- Morikawa S., Mabuchi Y., Kubota Y., Nagai Y., Niibe K., Hiratsu E., Suzuki S., Miyauchi-Hara C., Nagoshi N., Sunabori T., Shimmura S., Miyawaki A., Nakagawa T., Suda T., Okano H., Matsuzaki Y 2009) **Prospective identification, isolation, and systemic transplantation of multipotent mesenchymal stem cells in murine bone marrow** *J. Exp. Med* **206**:2483–2496
- Nusspaumer G., Jaiswal S., Barbero A., Reinhardt R., Ishay Ronen D., Haumer A., Lufkin T., Martin I., Zeller R 2017) **Ontogenic identification and analysis of mesenchymal stromal cell populations during mouse limb and long bone development** *Stem Cell Rep* **9**:1124–1138
- Pliner HA., Packer JS., McFaline-Figueroa JL., Cusanovich DA., Daza RM., Aghamirzaie D., Srivatsan S., Qiu X., Jackson D., Minkina A., Adey AC., Steemers FJ., Shendure J., Trapnell C 2018) **Cicero Predicts cis-Regulatory DNA Interactions from Single-Cell Chromatin Accessibility Data** *Mol. Cell* **71**:858–871
- Ruzzini L., Abbruzzese F., Rainer A., Longo UG., Trombetta M., Maffulli N., Denaro V 2014) **Characterization of age-related changes of tendon stem cells from adult human tendons** *Knee Surg Sports Traumatol Arthrosc* **22**:2856–66
- Satpathy AT., Granja JM., Yost KE., Qi Y., Meschi F., McDermott GP., Olsen BN., Mumbach MR., Pierce SE., Corces MR., Shah P., Bell JC., Jhutti D., Nemec CM., Wang J., Wang L., Yin Y., Giresi PG., Chang ALS., Zheng GXY., Greenleaf WJ., Chang HY 2019) **Massively parallel single-cell**

chromatin landscapes of human immune cell development and intratumoral T cell exhaustion *Nat Biotechnol* **37**:925–936

Sidney LE., Branch MJ., Dunphy SE., Dua HS., Hopkinson A 2014) **Concise review: evidence for CD34 as a common marker for diverse progenitors** *Stem Cells* **32**:1380–9

Somerville JM., Aspden RM., Armour KE., Armour KJ., Reid DM 2004) **Growth of C57BL/6 mice and the material and mechanical properties of cortical bone from the tibia** *Calcif Tissue Int* **74**:469–75

Staverosky JA., Pryce BA., Watson SS., Schweitzer R 2019) **Tubulin polymerization-promoting protein family member 3, Tppp3, is a specific marker of the differentiating tendon sheath and synovial joints** *Dev Dyn.* **238**:685–92

Steinmann S., Pfeifer CG., Brochhausen C., Docheva D 2020) **Spectrum of Tendon Pathologies: Triggers, Trails and End-State** *Int J Mol Sci* **21**:844

Still C, Chang WT., Sherman SL., Sochacki KR., Dragoo JL., Qi LS. 2021) **Single-cell transcriptomic profiling reveals distinct mechanical responses between normal and diseased tendon progenitor cells** *Cell Rep Med.* **2**:100343

Stuart T., Hanna J., Gutruf P 2021) **Single-cell chromatin state analysis with Signac** *Nat Methods* **18**:1333–1341

Sung JH., Yang HM., Park JB., Choi GS., Joh JW., Kwon CH., Chun JM., Lee SK., Kim SJ 2008) **Isolation and characterization of mouse mesenchymal stem cells** *Transplant Proc* **40**:2649–54

Tachibana N., Chijimatsu R., Okada H., Oichi T., Taniguchi Y., Maenohara Y., Miyahara J., Ishikura H., Iwanaga Y., Arino Y., Nagata K., Nakamoto H., Kato S., Doi T., Matsubayashi Y., Oshima Y., Terashima A., Omata Y., Yano F., Maeda S., Ikegawa S., Seki M., Suzuki Y., Tanaka S., Saito T 2022) **RSPO2 defines a distinct undifferentiated progenitor in the tendon/ligament and suppresses ectopic ossification** *Sci Adv* **8**:eabn2138

Takakura N., Yoshida H., Ogura Y., Kataoka H., Nishikawa S., Nishikawa S 1997) **PDGFR alpha expression during mouse embryogenesis: immunolocalization analyzed by whole-mount immunohistostaining using the monoclonal anti-mouse PDGFR alpha antibody APA5 J.** *Histochem. Cytochem* **45**:883–893

Tang JB., Wu YF., Cao Y., Chen CH., Zhou YL., Avanesian B., Shimada M., Wang XT., Liu PY 2016) **Basic FGF or VEGF gene therapy corrects insufficiency in the intrinsic healing capacity of tendons** *Sci Rep* **6**:20643

Tempfer H., Wagner A., Gehwolf R., Lehner C., Tauber M., Resch H., Bauer HC 2009) **Perivascular cells of the supraspinatus tendon express both tendon- and stem cell-related markers** *Histochem Cell Biol* **131**:733–41

Thomopoulos S., Parks WC., Rifkin DB., Derwin KA 2015) **Mechanisms of tendon injury and repair** *J Orthop Res* **33**:832–9

Trapnell C., Cacchiarelli D., Grimsby J., Pokharel P., Li S., Morse M., Lennon NJ., Livak KJ., Mikkelsen TS., Rinn JL. 2014) **The dynamics and regulators of cell fate decisions are revealed by pseudotemporal ordering of single cells** *Nat Biotechnol* **32**:381–386

Tsutsumi H., Kurimoto R., Nakamichi R., Chiba T., Matsushima T., Fujii Y., Sanada R., Kato T., Shishido K., Sakamaki Y., Kimura T., Kishida A., Asahara H 2022) **Generation of a tendon-like tissue from human iPS cells** *J Tissue Eng* **13**:20417314221074018

Walia B., Huang AH 2019) **Tendon stem progenitor cells: Understanding the biology to inform therapeutic strategies for tendon repair** *J Orthop Res* **37**:1270–1280

Wang B., Liu W., Zhang Y., Jiang Y., Zhang WJ., Zhou G., Cui L., Cao Y 2008) **Engineering of extensor tendon complex by an ex vivo approach** *Biomaterials* **29**:2954–61

Yin Z., Hu JJ., Yang L., Zheng ZF., An CR., Wu BB., Zhang C., Shen WL., Liu HH., Chen JL., Heng BC., Guo GJ., Chen X., Ouyang HW 2016) **Single-cell analysis reveals a nestin + tendon stem/progenitor cell population with strong tenogenic potentiality** *Sci Adv* **2**:e1600874

Yu G., Wang LG., Han Y., He QY 2012) **clusterProfiler: an R package for comparing biological themes among gene clusters** *Omics* **16**:284–7

Yu G., Wang LG., He QY 2015) **ChIPseeker: an R/Bioconductor package for ChIP peak annotation, comparison and visualization** *Bioinformatics* **31**:2382–3

Zhang C., Zhu J., Zhou Y., Thampatty BP., Wang JH 2019) **Tendon Stem/Progenitor Cells and Their Interactions with Extracellular Matrix and Mechanical Loading** *Stem Cells Int* **2019**:3674647

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Reviewer #1 (Public review):

This study is focused on identifying unique, innovative surface markers for mature Achilles tendons by combining the latest multi-omics approaches and in vitro evaluation, which would address the knowledge gap of the controversial identity of TPSCs with unspecific surface markers. The use of multi-omics technologies, in vivo characterization, in vitro standard assays of stem cells, and in vitro tissue formation is a strength of this work and could be applied for other stem cell quantification in musculoskeletal research. The evaluation and identification of Cd55 and Cd248 in TPSCs have not been conducted in tendons, which is considered innovative. Additionally, the study provided solid sequencing data to confirm co-expressions of Cd55 and Cd248 with other well-described surface markers such as Ly6a, Tpp3, Pdgfra, and Cd34. Generally, the data shown in the manuscript support the claims that the identified surface antigens mark TPSCs in juvenile tendons.

However, there are missing links between scientific questions aimed to be addressed in Introduction and Methodology/Results. If the study focuses on unsatisfactory healing responses of mature tendons and understanding of mature TPSCs, at least mature Achilles tendons from more than 12-week-old mice and their comparison with tendons from

juvenile/neonatal mice should be conducted. However, either 2-week or 6-week-old mice, used for characterization here, are not skeletally mature. Additionally, there is a lack of complete comparison of TPSCs between 2-week and 6-week-old mice in the transcriptional and epigenetic levels.

In order to distinguish TPSCs and characterize their epigenetic activities, the authors used scRNA-seq, snRNA-seq, and snATAC-seq approaches. The integration, analysis, and comparison of sequencing data across assays and/or time points is confusing and incomplete. For example, it should be more comprehensive to integrate both scRNA-seq and snRNA-seq data (if not, why both assays were used for Achilles tendons of both 2-week and 6-week timepoints). snRNA-seq and snATAC-seq data of 6-week-old mice were separately analyzed. No comparison of difference and similarity of TPSCs of 2-week and 6-week-old mice was conducted.

Given the goal of this work to identify specific TPSC markers, the specificity of Cd55 and Cd248 for TPSCs is not clear. First, based on the data shown here, Cd55 and Cd248 mark the same cell population which is identified by Ly6a, TPPP3, and Pdgfra. Although, for instance, Cd34 is expressed by other tissues as discussed here, no data/evidence is provided by this work showing that Cd55 and Cd248 are not expressed by other musculoskeletal tissues/cells. Second, the immunostaining of Cd55 and Cd248 doesn't support their specificity. What is the advantage of using Cd55 and Cd248 for TPSCs compared to using other markers?

<https://doi.org/10.7554/eLife.104768.1.sa2>

Reviewer #2 (Public review):

Summary:

The molecular signature of tendon stem cells is not fully identified. The endogenous location of tendon stem cells within the native tendon is also not fully elucidated. Several molecular markers have been identified to isolate tendon stem cells but they lack tendon specificity. Using the declining tendon repair capacity of mature mice, the authors compared the transcriptome landscape and activity of juvenile (2 weeks) and mature (6 weeks) tendon cells of mouse Achilles tendons and identified CD55 and CD248 as novel surface markers for tendon stem cells. CD55+ CD248+ FACS-sorted cells display a preferential tendency to differentiate into tendon cells compared to CD55neg CD248neg cells.

Strengths:

The authors generated a lot of data on juvenile and mature Achilles tendons, using scRNAseq, snRNAseq, and ATACseq strategies. This constitutes a resource dataset.

Weaknesses:

The analyses and validation of identified genes are not complete and could be pushed further. The endogenous expression of newly identified genes in native tendons would be informative. The comparison of scRNAseq and snRNAseq datasets for tendon cell populations would strengthen the identification of tendon cell populations.

<https://doi.org/10.7554/eLife.104768.1.sa1>

Reviewer #3 (Public review):

Summary:

In their report, Tsutsumi et al., use single nucleus transcriptional and chromatin accessibility analyses of mouse achilles tendon in an attempt to uncover new markers of tendon stem/progenitor cells. They propose CD55 and CD248 as novel markers of tendon stem/progenitor cells.

Strengths:

This is an interesting and important research area. The paper is overall well written.

Weaknesses:

Major problems:

- (1) It is not clear what tissue exactly is being analyzed. The authors build a story on tendons, but there is little description of the dissection. The authors claim to detect MTJ and cartilage cells, but not bone or muscle cells. The tendon sheath is known to express CD55, so the population of "progenitors" may not be of tendon origin.
- (2) Cluster annotations are seemingly done with a single gene. Names are given to cells without functional or spatial validation. For example, MTJ cells are annotated based on *Postn*, but it is never shown that *Postn* is only expressed at the MTJ, and not in other anatomical locations in the tendon.
- (3) The authors compare their data to public data based on interrogating single genes in their dataset. It is now standard practice to integrate datasets (eg, using harmony), or at a minimum using gene signatures built into Seurat (eg *AddModuleScore*).
- (4) Progenitor populations (SP1, SP2). The authors claim these are progenitors but show very clearly that they express macrophage genes. What are they, macrophages or fibroblasts?
- (5) All omics analysis is done on single data points (from many mice pooled). The authors make many claims on *n*=1 per group for readouts dependent on sample number (eg frequency of clusters).
- (6) The scRNAseq atlas in Figure 1 is made by analyzing 2W and 6W tendons at the same time. The snRNAseq and ATACseq atlas are built first on 2W data, after which the 6W data is compared. Why use the 2W data as a reference? Why not analyze the two-time points together as done with the scRNAseq?
- (7) Figure 5: The authors should show the gating strategy for FACS. Were non-fibroblasts excluded (eg, immune cells, endothelia...etc). Was a dead cell marker used? If not, it is not surprising that fibroblasts form colonies and express fibroblast genes when compared to CD55-CD248- immune cells, dead cells, or debris. Can control genes such as *Ptprc* or *Pecam1* be tested to rule out contamination with other cell types?

Minor problems:

- (1) Report the important tissue processing details: type of collagenase used. Viability before loading into 10x machine.

<https://doi.org/10.7554/eLife.104768.1.sa0>