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## The inflammatory response of the supraspinatus muscle in rotator cuff tear conditions.

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Running title: Muscle inflammation and rotator cuff

Conflicts of interest: None

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## Ethics

Ethical approval was granted by The Regional Committees on Health Research Ethics for Southern Denmark, J. No. S-20160037, and the study reported to The Danish Data Protection Agency (16/9714). The project was approved by the Orthopedic Research Board, Odense University Hospital.

1 2 The inflammatory response of the supraspinatus muscle in rotator cuff tear conditions. Running title: Muscle inflammation and rotator cuff

3

## 4 Abstract

Background: Rotator cuff (RC) disorders involve a spectrum of shoulder conditions from early
tendinopathy to full-thickness tears leading to impaired shoulder function and pain. The pathology
of RC disorder is, nonetheless, still largely unknown. It is our hypothesis that supraspinatus (SS)
tendon tear leads to sustained inflammatory changes of the SS muscle along with fatty infiltration
and muscle degeneration, which are threshold markers for poor RC muscle function. The aim of this
study was to determine the extent of this muscle inflammation in conjunction with lipid
accumulation and fibrosis in RC tear conditions.

12 **Methods:**We used proteomics, histology, electrochemiluminescence immunoassay, and qPCR 13 analyses to evaluate inflammatory and degenerative markers and fatty infiltration in biopsies from 14 22 patients undergoing surgery with repair of a full- thickness supraspinatus (SS) tendon tear. 15 Results: Bioinformatic analysis showed that proteins involved in innate immunity, extracellular 16 matrix organization, and lipid metabolism were among the most upregulated whereas mitochondrial 17 electronic transport chain along with muscle fiber function were among the most downregulated. 18 Histological analysis confirmed changes in muscle fiber organization and the presence of 19 inflammation and fatty infiltration. Inflammation appeared to be driven by a high number of 20 infiltrating macrophages, accompanied by elevated matrix metalloprotease levels and changes in 21 transforming growth factor- $\beta$  and cytokine levels in the SS compared to the deltoid muscle. 22 Conclusions: We demonstrated massive SS muscle inflammation after tendon tear combined with 23 fatty infiltration and degeneration. The regulation of tissue repair is thus extremely complex, and it 24 may have opposite effects at different time points of healing. Inhibition or stimulation of muscle 25 inflammation may be a potential target to enhance outcome of the repaired torn RC.

26 Level of evidence: Basic Science Study; Histology, Molcular and Cell Biology

27 Keywords: Shoulder disorder, muscle damage, proteomics, protein changes, extracellular matrix

- 28 degeneration, fatty infiltration
- 29

## 30 Introduction

31 Rotator cuff (RC) lesions are some of the most common shoulder conditions in humans and can 32 lead to weakness, pain, and limited/reduced mobility. The prevalence of RC tears is age-dependent, and both partial and full-thickness RC tears increase markedly after 50 years of age <sup>48</sup>. The etiology 33 34 of RC diseases is multifactorial with frequent involvement of the supraspinatus (SS) tendon<sup>5</sup>. Fullthickness SS tears do not heal spontaneously, and surgically repaired RC tears tend to heal poorly 35 <sup>15; 33</sup>. A recent Cochrane review questioned whether repair of RC tears provides meaningful benefit 36 to patients with symptomatic RC tears<sup>24</sup>. There is, therefore, a pressing need to better understand 37 the pathophysiology behind RC tear conditions in order to improve results after surgical repair of 38 39 RC tendon tears.

In full-thickness RC tears, increased numbers of immune cells have been demonstrated in the synovial tissue adjacent to the SS tendon <sup>1</sup> and an increase in tear size correlated with a greater proinflammatory response in the synovium <sup>6; 45</sup>. Recent data suggest that the RC muscles also become inflamed in the presence of an RC tear <sup>14</sup>, and results from experimental models suggest that acute inflammation plays a detrimental role in the onset of chronic muscle damage following RC tears <sup>16; <sup>29</sup>. It is also generally agreed that chronic RC tendon lesion leads to degenerative muscle changes in the form of fatty infiltration and fibrosis <sup>10</sup>.</sup>

47 Several animal studies have provided evidence of significant increases in inflammatory cytokines,

48 growth factors, and matrix metalloproteases (MMPs), indicating muscle inflammation following

49 experimental RC tendon tear <sup>16; 29</sup>. Changes in the biology of human RC muscles in tear conditions

remain poorly defined at the cellular level, however, and RC muscle as a target for inflammation
following RC tear is only sparsely understood <sup>14; 23</sup>.

52 The aim of this study was to provide a more robust understanding of the inflammatory environment

53 of human SS muscle in RC tear conditions. It is our hypothesis that inflammatory conditions

54 precede disturbances of the muscle architecture, and eventually lead to RC muscle fatty infiltration

and degeneration. To investigate this, we applied quantitative proteomics followed by histological,

56 multiplex chemiluminescence, and qPCR analyses of inflammation in SS and deltoid muscle

57 biopsies that were harvested from patients undergoing surgery for RC tears.

58

## 59 Materials and methods

## 60 Patient cohort

61 Patients (n=22) with a relevant shoulder trauma and clinical signs of an RC lesion were recruited

62 (Supplementary Table 1). Median age of lesions was 3.3 months (IQR 2-14 months). Patients

63 underwent preoperative magnetic resonance imaging scan revealing SS tendon tear, and all tears

were confirmed at surgery. Informed written consent was obtained from all patients. The workflowof the project is presented in Figure 1.

66

### 67 Human tissue

The RC and musculotendinous junction of the SS muscle were gently débrided from fascia and bursal tissue using a blunt shaver. SS tendon and muscle biopsies were harvested from the edge of the tendon and approximately one centimeter medial to the tendon, respectively, under direct visualization from the arthroscope. <u>C</u>omparative biopsies were taken from assumed healthy, ipsilateral deltoid muscles. Biopsies were snap-frozen on dry ice and stored at -80°C or fixed in 10% neutral buffered formalin and embedded in paraffin.

7	Λ.
/	т

## 75 Blood samples

76 Blood samples were obtained in EDTA coated test tubes immediately prior to surgery to estimate

77 preoperative peripheral inflammation. Hemoglobin, C-reactive protein and white blood cell counts

78 were analyzed. The patient cohort was also tested for presence of rheumatic factors (anti-nuclear,

anti-cyclic citrullinated, and mitochondrial antibodies and rheumafactor).

80

## 81 Histology and Immunohistochemistry

82 SS or deltoid muscle tissue was sectioned into 2µm thick sections on a microtome.

83 Immunohistochemistry for CD68 was performed on the OMNIS platform (Dako/Agilent, Denmark)

84 using mouse anti-CD68 (1:50, Clone PG-M1) antibody and EnVision<sup>TM</sup> FLEX as detection system,

85 while immunohistochemistry for FOXP3, CD3, and adipophilin was performed on a BenchMark

86 Ultra immunostainer (Ventana Medical Systems, AZ, USA) using mouse anti-FOXP3 (1:40, clone

87 236A/E7), rabbit anti-CD3 ("ready-to-use", clone 2GV6), and mouse anti-adipophilin (1:50, clone

AP125) antibodies and the OptiView-DAB detection system. Parallel sections were stained with

89 hematoxylin and eosin (HE).

90 Slides were scanned on a NanoZoomer 2.0 RS scanner (Hamamatsu Photonics, Visiopharm,

91 Denmark). To produce merged images, NDP view (Hamamatsu Photonics, version 2.6.17) was

92 applied to find identical regions on neighboring cells stained for FOXP3 and CD3, respectively.

93 The images were processed and merged using Photoshop C6 (Adobe Systems, CA, USA) and

94 ImageJ/Fiji. The area with muscle tissue in each biopsy was determined using the freehand region

95 function and the number of CD3 and FOXP3 within this area was manually counted at 20x

96 magnification.

97

#### 98 **Proteomics**

## 99 Homogenization of tissue biopsies

100 Approximately 1mm<sup>3</sup> was cut from the frozen tendon and muscle biopsies and was homogenized by

101 bead beating using ss 0.9-2.0mm beads (Bullet blender Gold, Next Advance Inc, USA) in reducing

- 102 lysis buffer (5% sodium deoxycholate (SDC; Sigma Aldrich, USA), protease inhibitors (cOmplete,
- 103 Roche) in 50mM TEAB (Sigma Aldrich). Protein concentration in lysates was determined by

104 protein A280 and lysates stored at -80°C (DeNoviX, USA).

105 Samples were further processed using an optimized SDC filter-aided sample preparation protein

106 digestion (SDC-FASP) essentially as <sup>4</sup> using 100µg protein starting material in 10kDa molecular

107 weight cutoff spin-filters (Merch Millipore, Singapore). The lysates were reduced (10mM TCEP)

and alkylated (50mM Chloroacetamide) each for 15min in digestion buffer (5% SDC in 50mM

109 TEAB; Sigma-Aldrich). Overnight trypsin digestion at 37°C was performed by addition of 200µl

110 0.5% digestion buffer containing Trypsin 1:50 (v:v; Pierce, USA). Peptides were extracted by

111 acidification and phase-separation, where 3:1 (v/v) ethyl acetate of sample volume was added and

acidified by adding Trifluoroacetic acid (Sigma-Aldrich) to a final concentration of 1% (pH<2)

followed by centrifigation. The collected lower aqueous phase containing the peptides was dried in

a vacuum centrifuge and dissolved in 0.1M TEAB. Peptide concentration in lysates was determined

115 by protein A280 and lysates stored at -80°C (DeNoviX, USA).

116 *iTRAQ* labeling

A total of 5µg of each sample was labeled with a 4-plex iTRAQ kit (AB Sciex, USA) according to
manufacturer's instructions. Briefly, samples were re-dissolved to total volume of 25µL 0.1M
TEAB, pH 8.5 while 290µL 96% ethanol were added to iTRAQ reagents. Next, 50µL of the
iTRAQ reagents was transferred to the samples, which were then labelled, mixed after 1h of

- 121 incubation at room temperature dried, and resuspended in 2% acetonitrile (AcN), 0.1% TFA
- 122 (Sigma-Aldrich).
- 123 *iTRAQ* sample analysis

124 Samples were analyzed per UPLC-TandemMS in technical duplicates. Labeled peptides were

- separated by a nanoUPLC system (Thermo Scientific, USA) coupled online to a Q Exactive HF MS
- 126 (Thermo Scientific) using a reverse phase C18 trapping column setup with 75cm main column
- 127 (Thermo Scientific) with loading in 2% solvent B (0.1% FA in AcN) gradient and separated by
- 128 176min gradient from 11%B to 30%B with a constant flow rate at 250nL/min. MS was operated in
- 129 positive mode using Top10 data-dependent acquisition (MS1 m/z 375-1,500 at R 120,000) and
- tandems sequencing using fixed m/z range at 110 and a MS2 resolution of 15,000.
- 131 Database searches

134

132 Raw data were processed using Proteome Discoverer v2.3.0.523 (Thermo Scientific). Sequest HT

133 was set as the search engine against the reviewed Uniprot Homo sapiens reference protein database

(09/2017). iTRAQ 4-plex labeling of N-terminal and lysine and carbamidomethylation (C) were set

- 135 as fixed modifications while oxidation (M), deamidation (N/Q) and protein N-terminal acetylation
- 136 were included as variable modifications. Precursor mass tolerance and fragment mass tolerance
- 137 were set at 10ppm and 0.05Da, respectively. PSMs were filtered using percolator with a strict false
- discovery rate (FDR) of 1% and a relaxed FDR of 5%. Unique and razor peptides were used for
- 139 quantification, and iTRAQ channels were normalized to total peptide amount. Master proteins were
- 140 filtered for high protein FDR confidence. MS data have been deposited to the ProteomeXchange
- 141 Consortium via the PRIDE <sup>39</sup> partner repository with the dataset identifier PXD014037.
- 142 Bioinformatics analyses and functional annotation of regulated proteins
- 143 Normalized abundances were used for further analyses of proteins identified with two or more
- unique peptides. Data distribution was assessed with Perseus v.1.5.3.2 software and differentially

145	regulated features were selected using t-test with a post-hoc background-based adjusted p-
146	value<0.05 <sup>35</sup> . Venny 2.1 (http://bioinfogp.cnb.csic.es/tools/venny) was used to compare the
147	regulated proteins among the different comparisons. ToppGene Suite <sup>8</sup> was used for functional
148	enrichment of regulated proteins according to Gene Ontology (GO) terms and pathway analysis.
149	Enriched lists were further accessed by String app on Cytoscape v3.6.1 <sup>44</sup> .
150	
151	Reverse transcription quantitative PCR (RT-qPCR) analysis of FOXP3, MYOG, and MMP13
152	in SS and deltoid muscle tissue
153	RNA extraction
154	Muscle biopsies (n=18/group) were isolated using TRIzol <sup>®</sup> Reagent. Phase separation was
155	performed using chloroform and isopropyl alcohol was used to precipitate RNA. The RNA
156	concentrations and purities were determined using a Nanodrop Spectrophotometer (Thermo
157	Scientific).
158	cDNA synthesis
159	RNA samples were diluted to obtain a concentration of $250 ng/\mu L$ , and reverse transcription was
160	performed using an Applied Biosystem kit according to the manufacturer's instructions. The
161	synthesis was performed using an MJ Research PTC-225 Gradient Thermal Cycler (Marshall
162	Scientific). cDNA samples were diluted to lower the concentrations to $\sim 50 \text{ng}/\mu L$ .
163	RT-qPCR
164	RT-qPCR was performed using a CFX Connect Real-Time PCR Detection System (Bio-Rad) and
165	analyzed using SYBR green. Samples were run against standard curves generated from serial
166	dilutions from a pool of all samples. Values were normalized to ACTB ( $\beta$ -actin) as the reference
167	gene and calibrated to a pool of cDNA obtained from one pectoralis and one subscapularis muscle
168	biopsy. Triplicates of all samples, standards, and negative controls were conducted. To ensure no

- sign of primer dimer formation or contamination, a no amplification control (NAC), a no template
- 170 control (NTC), and a no reverse transcriptase (NRT) were included as controls.
- 171 RT-qPCR cycling conditions were as follows: 10min at 95°C, followed by 40 cycles of denaturing
- 172 at 95°C for 15 seconds, 30 seconds at annealing temperature, and extension at 72°C at 30 seconds.
- 173 Primer sequences were: ACTB, sense 5'-GGCCACGGCTGCTTC-3' and anti-sense 5'-
- 174 GTTGGCGTACAGGTCTTTGC-3' (T<sub>a</sub> 52°C and T<sub>m</sub> 84°C), *FOXP3*, sense 5'-
- 175 CCCGGATGTGAGAAGGTCTT-3' and anti-sense 5'-TTCTCCTTCTCCAGCACCAG-3' (T<sub>a</sub>
- 176 57°C and T<sub>m</sub> 82°C), MYOG, sense 5'-GCCCTGATGCTAGGAAGCC-3' and anti-sense 5'-
- 177 CTGAATGAGGGCGTCCAGTC-3' (T<sub>a</sub> 70°C and T<sub>m</sub> 85°C), and *MMP13*, sense 5'-CGC CAG
- 178 ACA AAT GTG ACC CT-3' and anti-sense 5'-CAG GCG CCA GAA GAA TCT GT-3'\_(Ta 55°C
- and  $T_m$  77°C). Primer specificity was ensured by generation and evaluation of melting curves.
- 180 Primers were purchased from TAG Copenhagen.
- 181
- 182 Electrochemiluminescence analysis
- 183 *Protein purification*
- 184 Tendon and muscle samples were homogenized at 4°C in Mesoscale Lysis buffer containing
- 185 Phosphatase Inhibitor Cocktail 2 and 3 (Sigma-Aldrich) and Complete Mini EDTA-free Protease
- 186 Inhibitor (Roche). Protein content was measured by the bicinchoninic acid method using the
- 187 Thermo Scientific Micro BCA<sup>TM</sup> Protein assay Kit (Pierce Chemical Co)<sup>40</sup>.
- 188 Multiplex analysis
- 189 Protein concentrations in tendon and muscle samples were measured using an MSD human U-Plex
- 190 Biomarker Multip-Plex Kit, a U-PLEX human TGF-β Combo Kit, a human MMP 3-Plex Ultra-
- 191 Sensitive Kit, and human TNF-RI and TNF-RII Ultra-Sensitive Kits (all from Mesoscale), using the
- 192 MSD QuickPlex (SQ120) Plate Reader (Mesoscale) according to the manufacturer's instructions.

ICAM-1 and VCAM-1 analyses were performed on SS and deltoid muscle tissue using V-PLEX
Vascular Injury Panel 2 (Mesoscale). Samples were run in duplex and diluted 2- or 4-fold in Diluent
41 prior to measurement. Data were analyzed using MSD Discovery Workbench software. The
lower limit of detection was a calculated concentration based on a signal 2.5 standard deviations
(SD) above the blank (zero) calibrator and coefficient of variation (CV) values below 25% were
accepted.

199

## 200 Statistical analysis

201 To examine differences in protein expression between SS and deltoid muscle tissue, paired 202 Student's t-test was used. Correlation analyses between cytokine, MMPs, and growth factors versus 203 age of lesion used Spearman's test. To examine the correlation between the relative expression of 204 FOXP3, MYOG, and MMP13 in SS and deltoid muscles, a paired Wilcoxon test was carried out. 205 The outlier test ROUT was used to identify and remove outliers more than 2 SD from the dataset. 206 All statistical analyses were carried out using GraphPad Prism. P-values ≤0.05 were considered statistically significant. Data are presented as mean±SD or median (25, 75 interquartile range, 207 208 IQR).

209

210 **Results** 

## 211 Mass spectrometry-based proteomics analysis shows protein regulation upon RC lesion

Using quantitative mass spectrometry (MS) proteomics, a total of 2,463 proteins were identified, of which 1,895 had two or more unique peptides (Supplementary Table S2). Moreover, 417 quantified proteins were shared by all tissues of all patients and could be assessed by principal component analysis (PCA) (Supplementary Figure 1A), which showed a clear distribution pattern even in the absence of well-delimited clusters.

217	To better understand protein regulation underlying RC pathology, SS muscle protein expression
218	pattern was compared to deltoid as a non-RC shoulder muscle control. A total of 239 proteins were
219	regulated. Of these, 114 were more highly expressed in the SS muscle (Figure 2A, Supplementary
220	Table S3). Gene ontology analysis showed 'extracellular matrix organization' and 'neutrophil
221	degranulation' among the most enriched biological processes (Supplemental Figure 2, Table S4)
222	while 'degradation of extracellular matrix', 'innate immune system' and 'neutrophil degranulation'
223	were among the enriched pathways of upregulated proteins (Figure 2B, Supplementary Figure 2,
224	and Supplementary Table S4). Interestingly, 'mitochondrial cellular localization', and 'muscle
225	system process' were among the enriched annotations of downregulated proteins (Figure 2B,
226	Supplementary Figure 3, and Table S4). A detailed list of regulated proteins involved in the above-
227	mentioned processes, along with experimental ratio, are provided in Figure 2C-K.
228	To address the molecular response of different RC tissues, the comparison between SS muscle and
229	SS tendon showed 139 differentially expressed proteins (Supplementary Table S5). Pathways or
230	gene ontologies related to immune response or inflammatory processes were not enriched among
231	these regulated proteins.
232	In contrast, pointing towards a common molecular signature between SS muscle and SS tendon in
233	RC disease, 38 proteins were regulated in both tissues when compared to deltoid muscle
234	(Supplementary Figure 1B, Supplementary Tables S6 and S7). Several of these proteins were
235	related to catalytic activity, with mitochondrial protein complex being the main cellular component
236	(Supplementary Figure 1C).
237	
238	RC lesion results in atrophy and lipid accumulation
239	Staining with hematoxylin & eosin revealed muscle fiber changes already 11/2 months after tendon

240 lesion represented by internalization of nuclei and muscle fibers of varying size after the tendon tear

- 241 (Figure 3A). In addition, pathological infiltrating adipocytes were detected, as substantial fat
- infiltration was seen together with arrays of intracellular myonuclei and varying fiber size,
- 243 suggestive of degeneration (Figure 3B,C).
- Inflammatory changes appeared to be intensified from 1<sup>1</sup>/<sub>2</sub> months to 6 months with abundant
- stromal inflammatory phagocytic cells represented by the presence of CD68<sup>+</sup> macrophages (Figure
- 3D and 3E). At 24 months, degeneration was clearly seen with a substantial proportion of muscle
- cells replaced by fat cells indicative of muscle cell degeneration (Figure 3F). At 24 months, absent
- 248 or few CD68<sup>+</sup> positive cells indicated decreased inflammation. The deltoid muscle did not
- 249 demonstrate similar inflammatory and degenerative changes (Figure 3G-L).
- 250 Adipophilin/perilipin-2 immunohistochemistry for detection of lipids in myofibers demonstrated
- that adipophilin was localized to the surface of intra-cellular lipid droplets (Figure 4). The
- 252 expression varied both between patients (please compare Figure 4A and Figure 4C) and between
- 253 muscles from the same patient (please compare Figure 4A with Figure 4B). Regional differences
- within muscles were also seen (Figure 4), and some individual fibers presented increased lipid
- accumulation both as number and size of droplets.
- 256

## 257 RC tendon tear leads to changes in inflammatory mediators in SS muscle

None of the patients showed any signs of peripheral inflammation as mean blood leukocyte counts,
hemoglobin, and C-reactive protein values were within normal ranges and all patients were negative
for rheumatic factors. Given the findings of significant changes in the proteome of SS compared to
deltoid muscle after RC tear, we investigated changes in a variety of pro- and anti-inflammatory
cytokines, chemokines, receptors, and growth factors (Figure 5 and Table 1). We found lower
CCL19 levels (Figure 5A) but higher CXCL5 levels (Figure 5B) in the SS compared to the deltoid
muscle. IL-1β (Figure 5C), IL-6 (Figure 5D), IL-8 (Figure 5E), and IL-33 levels (Figure 5F) were

- higher in SS compared to deltoid muscle. IL-7 levels (Figure 5G), IL-15 (Figure 5H), IL-17A
- 266 (Figure 5I), and IFN-α2a levels (Figure 5J) were in SS compared to deltoid muscle. Despite
- 267 comparable levels of TNF (Figure 5K), TNFR1 (Figure 5L) and TNFR2 (Figure 5M) levels were
- 268 changed in the SS muscle compared to the deltoid after RC tendon tear. G-CSF levels were lower in
- 269 SS muscle compared to the deltoid (Figure 5N).
- 270 We observed no significant correlation between age of lesion and CCL19 (r=-0.05, p=1), CXCL5

271 (r=-0.26, p=0.31), IL-1β (r=-0.30, p=0.3), IL-6 (r=-0.48, p=0.07), IL-8 (r=-0.43, p=0.11), IL-33

272 (r=0.16, p=0.54), IL-7 (r=0.35, p=0.39), IL-15 (r=0.07, p=0.79), IL-17A (r=0.09, p=0.76), IFN-

- 273 α2a (r=0.39, p=0.3), TNF (r=-0.39, p=0.15), TNFR1 (r=-0.4, p=0.29), TNFR2 (r=-0.16, p=0.66),
- 274 or G-CSF (r=0.16, p=0.71).
- Finally, several cytokine levels were similar in SS and deltoid muscle (Table 1). The concentration
  of most cytokines, chemokines, growth factors, and MMPs was high in the SS tendon
- 277 (Supplementary Table 8).
- 278

## 279 The regenerative potential appears to be impaired in the SS muscle after RC tendon tear

280 IL-17 production characterizes pro-inflammatory T helper 17 lymphocytes (Th17) and innate immune cells <sup>11</sup>. Th17 has been shown to have opposing effects in the immune response from 281 regulatory T cells (Treg)<sup>49</sup>, which is important in muscle regeneration (reviewed in <sup>7</sup>) and whose 282 283 master gene is the transcription factor Forkhead box P3 (FOXP3). We found decreased IL-17A 284 levels in SS muscle compared to deltoid muscle (Figure 5G). Therefore, we next investigated 285 FOXP3 mRNA expression and found that FOXP3 mRNA levels were significantly lower in SS 286 muscle compared to deltoid muscle (Figure 6A). However, when we counted FOXP3<sup>+</sup> cells (Figure 287 6B) and CD3<sup>+</sup> T cells (Figure 6C) in SS and deltoid muscle tissue sections (Figure 6D), we did not observe any significant differences in the number of FOXP3<sup>+</sup> cells/mm<sup>2</sup> or CD3<sup>+</sup> T cells/mm<sup>2</sup>. 288

289	Approximately 7.5% of all T cells in the SS muscle were FOXP3 <sup>+</sup> Treg cells (overlay in Figure 6D,
290	upper panel), and approximately 13% of all T cells in the deltoid muscle were FOXP3 <sup>+</sup> Treg cells
291	(overlay in Figure 6D, middle panel). The number of T cells/mm <sup>2</sup> were, however, quite variable
292	(Figure 6D).
293	To investigate gene expression levels involved in myogenesis, we estimated MYOG expression and
294	found the relative mRNA levels to be significantly decreased in the SS compared to the deltoid
295	muscle (Figure 6E).
296	
297	Changes in matrix metalloproteinases and transforming growth factors after RC tendon tear
298	As gene ontology analysis showed changes in proteins involved in 'extracellular matrix
299	organization', we investigated changes in the levels of 4 matrix metalloproteinases (MMPs) known
300	to be involved in the degradation of the extracellular matrix $^{38}$ .
301	We did not observe any differences in MMP1 levels between SS and deltoid muscle (Figure 7A).
302	However, levels of MMP3 (that is known to degrade collagen types II-IV, IX, and X and to have
303	important regulatory functions such as activation of other MMPs) were significantly higher in SS
304	compared to deltoid muscle (Figure 7B). Furthermore, levels of MMP-9 (known to degrade
305	collagen fragments IV and V) were changed (Figure 7C) suggesting decreased MMP-9 levels in
306	deltoid compared to SS muscle.
307	MMP13 (known to degrade primarily collagen fragments II) mRNA gene expression was absent in
308	SS (0.005 $\pm$ 0.006, n=12) and deltoid (0.002 $\pm$ 0.002, n=16) muscle tissue, whereas <i>MMP13</i> mRNA
309	gene expression was present in SS tendon tissue $(4.95 \pm 5.33, n=4)$ in RC tear conditions.
310	A significant negative correlation was found between age of lesion and MMP9 levels (r=-0.61,
311	p=0.03). We found no significant correlation between age of lesion and MMP1 levels (r=0.05,
312	p=0.87) and a tendency of a correlation with MMP3 levels ((r=-0.46, p=0.08).

In line with previous findings in the SS enthesis (reviewed in  $^{22}$ ), we found that protein levels of 313 314 MMP-1 (known to specifically break down most subtypes of collagen, providing mechanical 315 strength to tissues) were higher in SS tendon than in muscle tissue (Supplementary Table 8). Also, 316 MMP-3 was high in the SS tendon (Supplementary Table 8). 317 We also investigated changes in transforming growth factors (TGF), known to be affected in SS enthesis following RC tear (reviewed in <sup>22</sup>) (Figure 7D-F). Levels of TGFB1 (Figure 7D) and 318 319 TGFB3 (Figure 7F) were higher in SS muscle, suggesting increased TGFB1 and TGFB3 levels in SS muscle compared to deltoid muscle. We found no correlation between age of lesion and TGF<sup>β</sup>1 320 321 (r=0.12, p=0.7), TGFβ2 (r=-0.07, p=0.83), or TGFβ3 (r=-0.32, p=0.34). 322

## 323 Discussion

In this study, we demonstrated massive SS muscle inflammation after tendon tear combined with
fatty infiltration and degeneration. Simultaneous changes in the innate immune response, cytokines,
and proteins related to extracellular matrix reorganization and mediation of fibrosis in the SS
musculature were also seen.

In line with previous experimental studies using mice <sup>28</sup> and rats <sup>12; 17; 18</sup> we observed high numbers of infiltrating monocytes/macrophages in SS muscle in early cases of tendon tear; this tendency ceased after 24 months, however. Inflammation was (initially) driven by high numbers of infiltrating CD68<sup>+</sup> macrophages, which are thought to be key sources of TGF- $\beta$ 1 linked to fibrosis in chronically injured muscle <sup>30</sup>.

333 Upregulated proteins in SS muscle compared to deltoid also showed 'neutrophil degranulation' 334 among the most enriched processes. Neutrophils have been identified as the main cells infiltrating 335 the muscle after injury <sup>46</sup>, and neutrophil-derived oxidants extended tissue damage in a rabbit model 336 of stretch skeletal muscle injury <sup>47</sup>. On the other hand, blocking cell infiltration compromised the 337 initial regenerative response, suggesting a role for neutrophils in muscle growth and repair by

338	removal of tissue debris and activation of satellite cells <sup>47</sup> . Moreover, a recent study has shown that
339	neutrophil-secreted proteases can also have an immunoregulatory role by activating IL-33 <sup>9</sup> . IL-33
340	has been shown to be produced by fibro-adipogenic progenitor (FAP) cells, which are uniformly
341	present in the interstitial space in skeletal muscle and respond to muscle damage $^{20}$ . Our
342	observation of higher IL-33 levels in SS muscle compared to deltoid muscle indicate an activation
343	of FAPs, which are the source of adipocytes in muscle fatty infiltration. Muscle Foxp3 <sup>+</sup> regulatory
344	T cells (Tregs) are characterized by high levels of expression of the IL-33 receptor, ST2, and are
345	known to potentiate regeneration in acute and chronic injury models <sup>7</sup> . Despite comparable numbers
346	of Foxp3 <sup>+</sup> Tregs/mm <sup>2</sup> , we saw a significant decrease in <i>FOXP3</i> mRNA levels in SS muscle
347	compared to deltoid muscle following RC tear, suggestive of repressed gene expression in Tregs
348	located in SS muscle. Many cytokines and factors can negatively regulate FOXP3 gene expression,
349	including IL-6, IL-7, TGF- $\beta$ , and G-CSF, <sup>21; 32; 42</sup> , all of which we observed to be different between
350	SS and deltoid muscle.
350 351	SS and deltoid muscle. The expression of IL-15 has been shown to inhibit fatty infiltration and facilitate muscle
350 351 352	SS and deltoid muscle. The expression of IL-15 has been shown to inhibit fatty infiltration and facilitate muscle regeneration through regulation of FAP cells <sup>23</sup> . In the present study, IL-15 levels were
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<ul> <li>350</li> <li>351</li> <li>352</li> <li>353</li> <li>354</li> <li>355</li> <li>356</li> <li>357</li> <li>358</li> </ul>	SS and deltoid muscle. The expression of IL-15 has been shown to inhibit fatty infiltration and facilitate muscle regeneration through regulation of FAP cells <sup>23</sup> . In the present study, IL-15 levels were significantly lower in the SS compared to the deltoid muscle, supporting our findings that adipocytes appear in the SS and that the regenerative potential appears to be reduced in SS muscle after RC tendon tear. IL-17 is a pro-inflammatory cytokine secreted by activated CD4 <sup>+</sup> T-helper cells (Th17), which are highly pro-inflammatory and induce severe autoimmunity (reviewed in <sup>27</sup> ). IL-17 levels are increased in early human tendinopathy, mediating inflammatory and tissue remodeling events <sup>34</sup> ,
<ul> <li>350</li> <li>351</li> <li>352</li> <li>353</li> <li>354</li> <li>355</li> <li>356</li> <li>357</li> <li>358</li> <li>359</li> </ul>	SS and deltoid muscle. The expression of IL-15 has been shown to inhibit fatty infiltration and facilitate muscle regeneration through regulation of FAP cells <sup>23</sup> . In the present study, IL-15 levels were significantly lower in the SS compared to the deltoid muscle, supporting our findings that adipocytes appear in the SS and that the regenerative potential appears to be reduced in SS muscle after RC tendon tear. IL-17 is a pro-inflammatory cytokine secreted by activated CD4 <sup>+</sup> T-helper cells (Th17), which are highly pro-inflammatory and induce severe autoimmunity (reviewed in <sup>27</sup> ). IL-17 levels are increased in early human tendinopathy, mediating inflammatory and tissue remodeling events <sup>34</sup> , and IL-17 inhibits myoblast differentiation <sup>26</sup> . In the present study, IL-17 levels were significantly
<ul> <li>350</li> <li>351</li> <li>352</li> <li>353</li> <li>354</li> <li>355</li> <li>356</li> <li>357</li> <li>358</li> <li>359</li> <li>360</li> </ul>	SS and deltoid muscle. The expression of IL-15 has been shown to inhibit fatty infiltration and facilitate muscle regeneration through regulation of FAP cells <sup>23</sup> . In the present study, IL-15 levels were significantly lower in the SS compared to the deltoid muscle, supporting our findings that adipocytes appear in the SS and that the regenerative potential appears to be reduced in SS muscle after RC tendon tear. IL-17 is a pro-inflammatory cytokine secreted by activated CD4 <sup>+</sup> T-helper cells (Th17), which are highly pro-inflammatory and induce severe autoimmunity (reviewed in <sup>27</sup> ). IL-17 levels are increased in early human tendinopathy, mediating inflammatory and tissue remodeling events <sup>34</sup> , and IL-17 inhibits myoblast differentiation <sup>26</sup> . In the present study, IL-17 levels were significantly lower in SS than in deltoid muscle, but the exact relevance of decreased IL-17 levels remains to be

362	Our gene ontology analysis showed changes in 'extracellular matrix organization' and 'degradation
363	of extracellular matrix' especially due to the upregulation of CMA1, COL5A3, CTSS, ELN, and
364	MMP19 in SS compared to deltoid muscle. To our knowledge, no one has investigated MMP levels
365	in human SS muscle under tear conditions. MMPs are a large group or proteolytic enzymes
366	responsible for tissue remodeling and degradation of extracellular matrix. In our study, MMP-3 and
367	MMP-9 levels were significantly increased in SS compared to deltoid muscle. MMP-3 is one of the
368	primary activators of MMP-9 from its inactive proenzyme form <sup>36</sup> . MMP-9 is produced by a variety
369	of cells, including fibroblasts. MMP-9 appears to be a regulatory factor in neutrophil migration
370	across the basement membrane <sup>13</sup> , and it also plays several important functions within neutrophil
371	action $^{27}$ such as degrading extracellular matrix, activation of IL-1 $\beta$ , and cleavage of several
372	chemokines <sup>37</sup> . In vitro studies have demonstrated that inhibition of MMP-9 reduced the levels of
373	active TGF- $\beta$ 1 and reduced several TGF- $\beta$ 1-driven responses such as fibroblast stimulation <sup>25</sup> . In
374	this context, MMP-9 appears to activate or stimulate the release of a number of cytokines and
375	growth factors, including TGF- $\beta$ 1 <sup>31</sup> , which we found to be elevated in SS compared to deltoid
376	muscle. MMP-9 activity positively correlated to skeletal muscle atrophy in immobilized rats <sup>41</sup> ,
377	supporting a role in muscle atrophy. This is in line with our present findings of a positive
378	correlation between MMP-9 levels and age of RC lesion. Altogether, this suggests that MMP-9
379	plays an important role in SS muscle remodeling.
380	While only a few studies have applied a large-scale proteomics approach to address RC

pathophysiology, these studies combined found several markers indicative of tissue remodeling and
 suggested an untapped potential for proteomics in tendon research (reviewed in <sup>43</sup>). A multi-omics
 methodology applied to a rat RC injury model to study myosteatosis identified disrupted

384 mitochondrial function as one of the underlying mechanisms of lipid accumulation in muscle fibers

<sup>19</sup>. In our study, we identified 32 downregulated mitochondrial proteins, 15 of which are associated

with mitochondrial electron transport chain. Mitochondrial dysfunction reduces energy production and the dysfunction of these organelles has been connected to the myosteatosis that is commonly reported following RC tendon injury <sup>3</sup>. Our finding of steatotic adipophilin positive muscle fibers and changes in lipid metabolism and mitochondrial function supports this connection.

390

391 This study has certain inherent limitations related to the variability of disease severity and duration 392 and the sample size, and the results are biased towards patients with RC tears who chose to undergo 393 surgery. The enrolled patients comprised four smokers and seven patients who received pain killers 394 on a daily basis. The adverse consequences of smoking and mild anti-inflammatory drug intake on 395 the inflammatory response was not assessed due to lack of statistical power. Another limitation 396 inherent in muscle biopsy studies is the difficulty of ensuring uniform biopsy procedures, which 397 are important to secure reproducibility and to account for regional variations in protein 398 composition. In this study, biopsies were obtained close to the musculotendinous junction of the SS 399 muscle in all patients using an all-arthroscopic approach from the bursal side, limiting possible 400 location-dependent variations. 401 The rationale for using the deltoid muscle for comparison could be challenged as it may be 402 asymptomatically affected. The ipsilateral deltoid has been used as a standard of reference in a

403 number of studies <sup>2; 13</sup>, also justifying the use of paired statistics and increasing the power of the
404 analyses.

405

## 406 Conclusions

This study demonstrated massive muscle changes after SS tendon tear characterized by high
numbers of inflammatory macrophages in lesions less than three months old and overall changes in
cytokine levels, MMP levels, and growth factors. Our proteome analysis demonstrated that proteins

410	involved in inflammation, extracellular matrix reorganization, and lipid metabolism were among the
411	most enriched. Knowing that massive inflammation with infiltration of immune cells into the RC
412	musculotendinous lesion disrupt normal muscle regeneration, this implies that intervention with
413	repair of the tendon lesion and concomitant target-specific adjuvant treatment of the inflammatory
414	state of the SS muscle could be key to improving RC muscle recovery.
415	
416	Figure legends
417	
418	Figure 1. Schematic workflow applied to the study of molecular pathways involved in the RC
419	lesion. Biopsies of supraspinatus tendon and supraspinatus and deltoid muscle biopsies from ten
420	patients undergoing surgery for partial or full-thickness RC were evaluated.
421	
422	Figure 2. Proteomics of muscle biopsies. (A) Differentially regulated protein pattern between SS
423	and deltoid muscles from patients with RC lesion (FDR 5%). Downregulated proteins are
424	represented in blue and upregulated in red. Dotted lines highlight proteins at least 2 times
425	overrepresented in each tissue. (B) Protein-protein network of SS vs deltoid muscle-regulated
426	proteins grouped according functional enrichment. Underrepresented proteins in SS muscle are
427	shown in blue while overrepresented are shown in red. (C-K) Levels of upregulated (red) or
428	downregulated (blue) proteins involved in Leukocyte mediated immunity (C), Immune effector
429	process (D), Immune system (E), Positive regulation of lipid metabolic processes (F), Metabolism
430	of lipids (G), Electron transport chain (H), Muscle structure development (J), Extracellular matrix
431	organization (I), and Striated muscle contraction (K) as measured by proteomics (n=10 muscles per
432	group). Abbreviations: ADAR, double-stranded RNA-specific adenosine deaminase; ADIPOQ,
433	adiponectin; ALDH5A1, succinate-semialdehyde dehydrogenase, mitochondrial; ANKRD1,
434	ankyrin repeat domain-containing protein 1; ANKRD2, ankyrin repeat domain-containing protein 2;
435	AP1B1, AP-complex subunit beta-1; APOA1, apolipoprotein A-1; APOA2, apolipoprotein A-2;
436	APOA4, apolipoprotein A-4; APOE, apolipoprotein E; CMA1, chymase; COL5A3, collagen alpha-
437	3(V) chain; COX6A1, cytochrome c oxidase subunit 6A1; COX6A2, cytochrome c oxidase subunit
438	6A2; COX6B1, cytochrome c oxidase subunit 6B1; CTSA, cathepsin A; CTSC, cathepsin C; CTSS,
439	cathepsin S; DNM1, dynamin-1; DNM2, dynamin-2; EEF1A2, elongation factor-1 alpha; ELN,

440 elastin; ENG, endoglin; F2, prothrombin; FABP4, fatty acid-binding protin; FASN, fatty acid 441 synthase; FBN1, fibrillin-1; FDXR, NADPH:adrenodoxin oxidoreductase, mitochondrial; FITM2, 442 fat storage-inducing transmembrane protein 2; GGH, gamma-glutamyl hydrolase; GLRX5, 443 glutaredoxin-related protein 5, mitochondrial; GNS, N-acetylglucosamine-6-sulfatase; GSR, 444 glutathione reductase, mitochondrial; HEXA, beta-hexosaminidase subunit alpha; HLA-A, HLA 445 class I histocompatibility antigen, A-31 alpha chain; HPGDS, hematopoietic prostaglandin D 446 synthase; HUWE1, E3 ubiquitin-protein ligase; ICAM1, intercellular adhesion molecule 2; 447 LGALS1, galectin-1; MFAP4, microfibril-associated glycoprotein 4; MMP19, matrix 448 metalloproteinase-19; MYH3, myosin-3; MYH7, myosin-7; MYL2, myosin light chain 2; MYL3, 449 myosin light chain 3; MYL6B, myosin light chain 6B; MYOZ2, myozenin-2; ND4, NADH-450 ubiquinone oxidoreductase chain 4; NDUFA4, cytochrome c oxidase subunit NDFUA4; NDUFA9, 451 NADH dehydrogenase [ubiquinone] 1 alpha subcomplex subunit 9, mitochondrial; NDUFAB1, 452 acyl carrier protein, mitochondrial; NDUFB3, NADH dehydrogenase [ubiquinone] 1 beta 453 subcomplex subunit 3; NDUFC2, NADH dehydrogenase [ubiquinone] 1 subunit C2; NDUFS5, 454 NADH dehydrogenase [ubiquinone] iron-sulfur protein 5; NDUFS7, NADH dehydrogenase 455 [ubiquinone] iron-sulfur protein 7, mitochondrial; NUP62, nuclear pore glycoprotein p62; OSTF1, 456 osteoclast-stimulating factor 1; PLIN1, perilipin-1; PON2, serum paraoxonase/arylesterase 2; PPT1, 457 palmitoyl-protein thioesterase 1; PRDX4, peroxiredoxin-4; PTPRC, receptor-type tyrosine-protein 458 phosphatase C; PYCARD, apoptosis-associated speck-like protein containing a CARD; SACM1L, 459 phosphatidylinositide phosphatase SAC1; SERPINB12, serpin B12; SLC25A1, tricarboxylate 460 transport protein, mitochondrial; SNRPA1, U2 small nuclear ribonucleoprotein A; SORBS1, sorbin 461 and SH3 domain-containing protein 1; TNNI1, troponin I, slow skeletal muscle; TNNT1, troponin 462 T, slow skeletal muscle; UCHL1, ubiquitinin carboxyl-terminal hydrolase isozyme L1; UQCR11, 463 cytochrome b-c1 complex subunit 10; VCAM1, vascular cell adhesion protein 1.

464

Figure 3. Histological analysis of SS muscle. (A-C) H&E-stained tissue sections of SS muscle biopsies from representative patients at 1½ months (A), 6 months (B), and 28 months (C) after RC tendon tear demonstrating the presence of muscle fibers with internal nuclei (arrows) and nuclear chains (arrow heads). (D-F) Immunohistochemical staining of SS muscle biopsies from representative patients at 2½ months (D), 6 months (E), and 28 months (F) demonstrating the presence of a high number of CD68<sup>+</sup> macrophages (arrows) within the first 6 months after RC

471 tendon tear. Please note the presence of massive fatty cell infiltration (\*, asterisks) between muscle

472 fibers already early after RC tendon tear. (G-I) H&E-stained tissue sections of deltoid biopsies from
473 representative patients at 1½ months (G), 6 months (H), and 24 months (I) after RC tendon tear. (J474 L) CD68 immunohistochemical staining of deltoid muscle biopsies biopsies from representative
475 patients at 1½ months (J), 6 months (K), and 24 months (L) after RC tendon tear. Scale bar: 100µm.
476
477 Figure 4. Immunohistochemical staining for adipophilin (A P) Adipophilin expression in SS.

- 477 Figure 4. Immunohistochemical staining for adipophilin. (A,B) Adipophilin expression in SS 478 (A) and deltoid (B) muscle fibers in a patient with a 6-month-old RC lesion, showing higher 479 adipophilin expression in the SS muscle compared to the deltoid. Adipophilin is seen as a granular 480 staining in the muscle fiber cytoplasm and due to its localization to the surface of lipid vacuoles, it 481 visualizes the distribution one of intracellular lipid. (C,D) Adipophilin expression in SS and deltoid 482 muscle fibers from patient with a >72-month-old RC tear (C) and another patient with a 7-month-483 old RC lesion (D). Please note that the distribution of adipophilin can be uneven with different 484 expression in neighboring fascicles in both the SS and deltoid muscles. Scale bar: 100µm.
- 485

486 Figure 5. Cytokine and TNF receptor protein expression in SS and deltoid muscle tissue in

487 **RC tear conditions.** (A-N) Electrochemiluminescence immunoassay analysis of CCL19 (A),

488 CXCL5 (B), IL-1β (C), IL-6 (D), IL-8 (E), IL-33 (F), IL-7 (G), IL-15 (H), IL-17A (I), INF-α2a (J),

489 TNF (K), TNFR1 (L), TNFR2 (M), and G-CSG (N) protein levels in SS and deltoid muscle

biopsies from patients with RC tendon tear. \*\*\*p<0.001, \*\*p<0.01. \*p<0.05, Student's t-test (n = 0.01) + 0.01.

491 22/group). Samples with CV values above 25% were excluded in individual analyses.

492

493 Figure 6. FOXP3 and MYOG mRNA expression is lower in SS muscle than in deltoid muscle

494 tissue in RC tear conditions. (A) FOXP3 mRNA levels in SS and deltoid muscle biopsies 495 demonstrated significantly lower expression levels in SS compared to deltoid muscle. \*\*p<0.01, 496 paired Student's t-test (n=18/group). Two outliers in the SS muscle and two outliers in the deltoid 497 muscle group were removed according to ROUT's outlier test. (B-C) The number of FOXP3<sup>+</sup> Treg cells/mm<sup>2</sup> (B) and the total number of CD3<sup>+</sup> T cells (C) were comparable between SS and deltoid 498 499 muscle in RC tear conditions. (D) Representative FOXP3 and CD3 immunohistochemically stained 500 tissue sections from SS and deltoid muscle demonstrating overlay between subsets of FOXP3<sup>+</sup> and 501 CD3<sup>+</sup> T cells, representing the presence of Treg cells (arrows) in both SS and deltoid muscle in RC 502 tear conditions. Scale bars: 50µm (top and middle panels) and 100µm (bottom panel). (E) MYOG

- 503 mRNA levels in SS and deltoid muscle biopsies demonstrated significantly lower expression levels
  504 in SS compared to deltoid muscle. \*\*p<0.01, paired Student's t-test (n=18/group).</li>
- 505
- 506 Figure 7. Matrix metalloprotease and transforming growth factor-β protein expression in SS

507 and deltoid muscle tissue under RC tear conditions. (A-F) Electrochemiluminescence

508 immunoassay analysis of MMP-1 (A), MMP-3 (B), MMP-9 (C), TGFβ1 (D), TGFβ2 (E), and

509 TGF $\beta$ 3 (F) protein levels in SS and deltoid muscle biopsies from patients with RC tendon tear.

- p<0.05, Student's test (n = 22/group). Samples with CV values above 25% were excluded in
- 511 individual analyses.
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## Table 1. Cytokine analysis in patients with RC tear.

	SS muscle (pg/mg)	Deltoid muscle (pg/mg)	P-value
IL-1α	2.44 ± 1.38 (n=13)	2.21 ± 1.72 (n=15)	0.83
IL-1Ra	46.83 ± 20.24 (n=14)	34.40 ± 12.94 (n=15)	0.33
IL-2	0.64 ± 0.47 (n=4)	0.41 ± 0.24 (n=5)	0.50
IL-2Ra	93.08 ± 56.28 (n=8)	99.11 ± 76.81 (n=7)	0.87
IL-4	0.04 ± 0.07 (n=6)	0.07 ± 0.08 (n=6)	0.53
IL-9	1.12 ± 0.72 (n=16)	1.10 ± 0.62 (n=17)	0.98
IL-12/IL-23p40	14.35 ± 29.01 (n=15)	14.64 ± 7.71 (n=18)	0.78
IL-12p70	0.23 ± 0.07 (n=6)	0.39 ± 0.32 (n=7)	0.55
IL-13	2.76 ± 2.25 (n=3)	4.35 ± 3.68 (n=3)	0.31
MIF	28,656 ± 18,419 (n=9)	26,617 ± 19,295 (n=8)	0.36
IFN-β	46.08 ± 27.05 (n=8)	34.05 ± 37.80 (n=7)	0.56
IFN-γ	2.23 ± 3.48 (n=5)	2.92 ± 2.11 (n=4)	0.95
FLT3L	14.29 ± 5.13 (n=5)	15.01 ± 8.78 (n=4)	0.93
TRAIL	54.41 ± 23.81 (n=17)	47.00 ± 17.70 (n=19)	0.50
CXCL1/GROα	2.63 ± 2.05 (n=9)	3.92 ± 3.37 (n=8)	0.16
CXCL10/IP-10	8.73 ± 4.00 (n=7)	8.69 ± 2.94 (n=9)	0.86
CXCL11/I-TAC	6.74 ± 2.02 (n=15)	7.78 ± 3.00 (n=19)	0.28
CCL2/MCP1	4.03 ± 2.38 (n=8)	4.83 ± 2.17 (n=9)	0.33
CCL3/MIP-1 α	4.58 ± 2.35 (n=5)	4.01 ± 4.05 (n=2)	-
CCL4/MIP-1β	5.17 ± 1.62 (n=9)	6.35 ± 1.54 (n=8)	0.29
CCL7/MCP-3	4.92 ± 2.59 (n=8)	4.03 ± 3.41 (N=6)	0.33
CCL8/MCP-2	1.52 ± 0.76 (n=8)	1.71 ± 1.11 (n=4)	0.48
CCL13/MCP4	15.64 ± 11.80 (n=8)	9.57 ± 7.33 (n=7)	0.24
CCL17/TARC	3.05 ± 1.01 (n=7)	3.40 ± 0.59 (n=8)	0.41
M-CSF	5.45 ± 3.46 (n=9)	6.27 ± 3.39 (n=6)	0.84
GM-CSF	0.13 ± 0.14 (n=3)	0.10 ± 0.11 (n=7)	0.33
LT-α	1.60 ± 0.79 (n=6)	1.67 ± 0.61 (n=9)	0.41
YKL-40/CHI3L1	121.40 ± 37.64 (n=5)	340.80 ± 274.40 (n=9)	0.18
VEGF-A	93.36 ± 31.19 (n=7)	134.40 ± 53.22 (n=9)	0.19

Paired Student's t-test



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