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9	The progression of infiltrating neurovascular features and chemokine production of the caudal
10	intervertebral disc following injury
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31	Keywords: intervertebral disc degeneration; inflammatory chemokines; intradiscal vascularization;
32	innervation; low back pain pathoanatomy

# 33 Abstract

34	The accessibility of the mouse caudal intervertebral disc (IVD) and its geometric semblance to the human
35	IVD makes it an attractive model for assessing IVD-specific responses in vivo. To effectively utilize this
36	model, the temporal trajectories of key pathoanatomical features, such as the production of inflammatory
37	chemokines, tissue disorganization, and neo- vessel and neurite infiltration, must be understood. This
38	study aims to define the progression of chemokine production and neurovascular invasion at 2-, 4-, and
39	12-weeks following a caudal IVD injury in 3-month-old female C57BL6/J mice. We measured IVD-
40	secreted chemokines and matrix metalloproteinases (MMPs) using multiplex ELISA, graded the
41	histopathological degeneration, and quantified the intradiscal infiltrating vessels (endomucin) and nerves
42	(protein-gene-product 9.5) using immunohistochemistry. Injury provoked the secretion of IL6, CCL2,
43	CCL12, CCL17, CCL20, CCL21, CCL22, CXCL2 and MMP2 proteins. Neurites propagated rapidly
44	within 2-weeks post-injury and remained relatively constant until 12-weeks. Peak vascular vessel length
45	occurred at 4-weeks post-injury and regressed by 12-weeks. These findings identified the temporal flux of
46	inflammatory chemokines and pain-associated pathoanatomy in a model of IVD degeneration using the
47	mouse caudal spine.

## 48 **1. Introduction**

Low back pain affects up to 85% of the population worldwide<sup>1,2</sup>, and intervertebral disc (IVD) 49 degeneration is a significant contributing factor.<sup>3</sup> The IVD is a cartilaginous soft tissue and is considered 50 51 avascular and aneural.<sup>4</sup> The functional spinal unit consists of the IVD and adjacent vertebral bodies where 52 the IVD provides resistance against compressive loads and shock absorbance for the musculoskeletal system. With degeneration, the IVD has compromised ability to perform these essential functions and can 53 become painful.<sup>3</sup> Some key hallmarks of IVD degeneration include disorganization of the IVD tissue, 54 secretion of inflammatory chemokines and growth factors<sup>5,6</sup>, and infiltration of blood vessels and 55 nerves.<sup>7,8</sup> 56

57 In addition to the structural collapse and the depletion of proteoglycan-rich matrix, other biological features in the degenerating IVD may be culprits to chronic pain, including the production of 58 inflammatory chemokines, expression of catabolic enzymes, and the invasion of neurites and vessels.<sup>9,10</sup> 59 Chemokines canonically recruit immune cells, which in turn secrete more chemokines that further 60 exacerbate the inflammatory state of the degenerating IVD<sup>11</sup>, and the immune cells can further accelerate 61 the breakdown of the extracellular matrix.<sup>12,13</sup> Chemokines produced by the IVD may also contribute to 62 63 neuron and vessel propagation around and into the outer annulus fibrosus which may mediate of IVD degeneration associated low back pain.<sup>14</sup> 64

Animal models are a common tool for studying IVD degeneration. Injury such as via mechanical 65 overload<sup>15–18</sup> and needle puncture<sup>19–33</sup> are used to induce degeneration of the IVD. Though the lumbar 66 spine is more clinically relevant as a site of pain generation compared to the caudal spine, the surgical 67 exposure required to access the lumbar IVD is traumatic, and the surrounding inflammation may 68 69 confound the IVD-specific responses. Puncture injury to the caudal IVD does not require surgical exposure and can be implemented with radiographic guidance.<sup>25</sup> Furthermore, the murine caudal spine 70 consists of 27 intervertebral discs, compared to just 5 in the lumbar spine,<sup>34</sup> and thus enable better control 71 72 of inter-animal variability by allowing comparisons of IVDs subjected to different treatment conditions

- vithin the same mouse. Therefore, the caudal spine may be more experimentally efficient for
- 74 investigating IVD-specific disease mechanisms.
- 75 To effectively leverage the advantages of the caudal model, it is crucial to define the progression
- of the inflammatory cascade and pain-related neurovascular features over time. Both neurites and vessels
- have been observed in aged mouse lumbar  $IVDs^{35}$  and in human degenerated  $IVDs^{8}$ , but the time course
- of how the caudal IVD will recapitulate these features is unclear. Therefore, the objective of this study is
- to define the temporal progression of neurites, vessels, and the IVD's production of chemokines during
- 80 injury-induced degeneration of the mouse caudal IVD.

#### 81 2. Materials and Methods

#### 82 2.1 Animal model

All animal procedures were performed with Washington University School of Medicine IACUC 83 84 approval. Three-month-old C57BL6/J female mice (N = 35) were used in this study. They were housed 85 under standard animal husbandry conditions (in a temperature-controlled  $[21 \square \pm \square 1^{\circ}C]$  room with normal 12-h light/dark cycles). Bilateral puncture with 30G needle of caudal (Coccygeal - Co) intervertebral 86 87 discs (IVD) was performed and adjacent IVDs were used as internal controls. Pre- and post-procedural Xray (Faxitron UltraFocus 100) was used to locate the IVDs of interest and confirm puncture. Co4/5 and 88 89 Co6/7 were injured with Co3/4 and Co7/8 acting as internal controls. A set of animals (n = 5) underwent a sham procedure to create a superficial injury where the skin and surrounding soft tissue was punctured. 90 91 Longitudinal assessment of pain behavior and locomotive performance was performed on a subset of 92 animals (Supplemental methods). Mice were euthanized at 2, 4 and 12 weeks (n = 9-10 per timepoint) 93 after injury; all sham mice were taken out to 12 weeks. Paired control and injured IVDs from bilateral 94 puncture mice were divided between OCT embedded histology (Co3/4 and Co4/5; n = 9-10 per timepoint), paraffin embedded histology (Co7/8 and Co6/7; n = 5 per timepoint) and organ culture (Co7/8 95 96 and Co6/7; n = 4-5 per timepoint). Sham control and punctured IVDs were divided between 97 immunohistochemistry (Co3/4 and Co4/5) and paraffin embedded histology (Co7/8 and Co6/7). The 98 lumbar dorsal root ganglions were also extracted from a subset of animals and underwent staining for 99 TRPV1 (Supplemental methods). All samples for histology were immediately fixed in 4% 100 paraformaldehyde at time of sacrifice for 24-48 hours. 101

102 2.2 Paraffin embedded histology

103 Spinal segments with Co7/8 and Co6/7 (n = 5 per timepoint) were embedded in paraffin and 10  $\mu$ m thick

104 sagittal sections were stained with Safranin-O against FAST green. IVDs were graded using a

standardized 35-point histopathology scale.<sup>36</sup> Three independent blinded individuals graded all

106 histological sections to consensus.

107 2.3 Quantification of Secreted Factors

108	Co7/8 and Co6/7 ( $n = 4-5$ each per timepoint) functional spinal units (FSUs) were immediately placed in
109	tissue culture media at time of sacrifice. Culture media consisted of 1:1 Dulbecco's modified Eagle's
110	medium: Nutrient mixture F-12 (DMEM:F12) supplemented with 20% fetal bovine serum and 1%
111	penicillin–streptomycin. FSUs were cultured for 6 days at $37^{\circ}$ C and 5% CO <sub>2</sub> with a complete media
112	change after 3 days. Day 6 media was collected and analyzed as described here.
113	
114	Chemokines were measured using the Luminex <sup>™</sup> 200 system (Luminex) conducted by Eve Technologies
115	Corp. using two separate kits, a 32-plex and a 12-plex assays (MilliporeSigma) to detect a total of 44
116	markers. The 32-plex included Eotaxin (CCL11), granulocyte colony-stimulating factor (GCSF),
117	granulocyte-macrophage colony-stimulating factor (GMCSF), IFN-γ, IL-1α, IL-1β, IL-2, IL-3, IL-4, IL-5,
118	IL-6, IL-7, IL-9, IL-10, IL-12 (p40), IL-12 (p70), IL-13, IL-15, IL-17, IP-10 (CXCL10), KC (CXCL1),
119	leukemia inhibitory factor (LIF), LIX (CXCL5), MCP-1 (CCL2), MCSF, MIG (CXCL9), MIP-1α
120	(CCL3), MIP-1 $\beta$ (CCL4), MIP-2 (CXCL2), RANTES (CCL5), TNF $\alpha$ , and VEGFA. The 12-plex
121	measured 6Ckine/Exodus2 (CCL21), Fractalkine (CX3CL1), IFN-\beta1, IL-11, IL-16, IL-20, MCP-5
122	(CCL12), MDC (CCL22), MIP-3α (CCL20), MIP-3β (CCL19), TARC (CCL17), and TIMP-1. Assay
123	sensitivities of these markers range from 0.3–30.6 pg/mL.
124	Matrix metalloproteases (MMPs) were quantified using a single 5-plex kit (MilliporeSigma). This
125	kit measured MMP-2, MMP-3, MMP-8, proMMP-9 and MMP-12. Assay sensitivities of these markers
126	range from 1.6 – 8.4 pg/mL. Individual analyte sensitivity values for all kits are available in the
127	MilliporeSigma MILLIPLEX® MAP protocol.
128	
129	2.4 Evaluation of intradiscal vascularization and innervation
130	Spinal segments including Co $3/4$ and Co $4/5$ IVDs (n = 9-10 per timepoint) were embedded in OCT and
131	sectioned along the sagittal plane at a 50 $\mu$ m thickness. Frozen sections were stained with anti-protein
132	gene product 9.5 (PGP9.5) and anti-endomucin (EMCN) against DAPI. PGP9.5 is a neuronal marker for

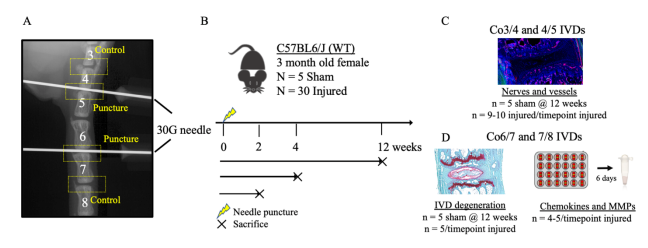
sensory and autonomic nerve fibers and EMCN is an endothelial cell marker. Visualization of PGP9.5
and EMCN was achieved with Alexa Fluor 488 (green) and Alexa Fluor 647 (red) antibodies,
respectively.

136 Three-dimensional image stacks were obtained via confocal fluorescence microscopy (DMi8, 137 Leica Microsystems) and a maximum intensity image of each 50  $\mu$ m section was generated for analysis. Nerves and vessels were semi-automatically traced using ImageJ 2.3.0 SNT plugin.<sup>37</sup> Individual structure 138 lengths were tabulated and total neurite and vessel length was calculated including both the posterior and 139 anterior sides of the IVD. The outer annulus fibrosus and immediately adjacent tissues were included as 140 the region of interest (ROI) for quantification. 141 142 2.6 Statistics 143 A paired two-way ANOVA was used to test for an effect of injury and week post-injury between the 144

experimental and control segments, at a significance level of 0.05 with a post hoc Turkey HSD (Prism

146 10.2.2, GraphPad). A paired t-test was used to test for an effect of the superficial injury on experimental

147 levels versus control segments in the sham-injured group only (12 weeks).

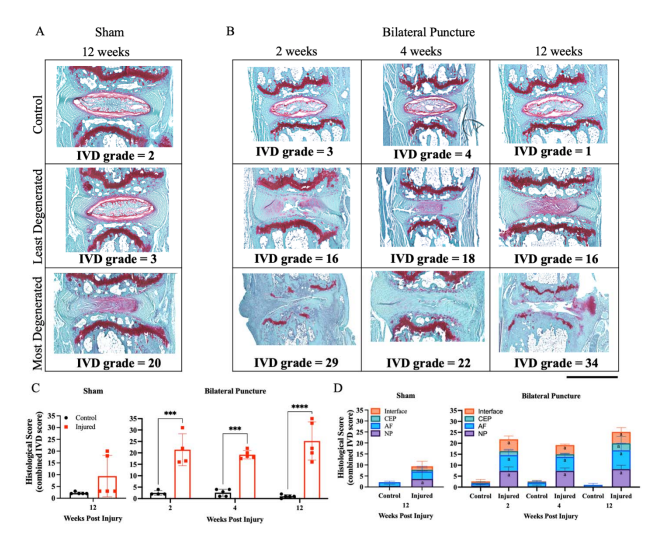


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- 149 Figure 1: <u>Bilateral caudal injury model in female C57BL6/J mice</u>. (A) Injury of Co4/5 and Co6/7 IVDs
- 150 were confirmed via X-ray. The Co3/4 and Co7/8 IVDs were used as internal controls within the same
- animal. (B) Bilateral puncture mice were sacrificed at 2, 4 and 12 weeks post injury while all Sham mice
- 152 were sacrificed at 12 weeks. Sham was a superficial injury limited to the skin and surrounding tissue. The
- internal controls (Co3/4 and 7/8) and injured (Co4/5 and 6/7) IVDs were divided between (C)
- 154 Immunohistochemistry for nerves (Green: protein gene product 9.5; PGP9.5) and vessels (Red:
- 155 endomucin; EMCN) stained against DAPI (Blue) and (D) Safranin-O against FAST Green staining for
- 156 IVD degeneration and tissue culture for IVD-specific chemokine and MMP production.

## 157 **Results**

- 158 3.1 Direct injury to the intervertebral disc causes rapid and sustained degeneration
- 159 Bilateral puncture of the caudal intervertebral disc (IVD) resulted in mild to severe IVD degeneration (Fig
- 160 2A). Complete collapse of the IVD was observed in the most severe cases. The control IVDs shown are
- 161 from the same animal that exhibited the most severe degeneration. Co6/7 and Co7/8 IVDs from sham and
- bilateral puncture mice were graded on a histopathologic scale for IVD degeneration and total IVD grade
- 163 was significantly increased in punctured IVDs compared to internal controls of injured (p<0.05, ANOVA)
- but no differences were detected in these levels in the sham mice (p=0.1, t-test; Fig. 2B). Multiple
- 165 compartments of the IVD showed degenerative changes, including the nucleus pulposus, annulus fibrosus
- and the interfaces at all timepoints after injury while cartilaginous endplates were only significantly
- 167 degenerated at 12 weeks following injury (Fig. 2C). No effect of injury was observed in any pain
- 168 behavior or locomotive assessments (Supplemental results).



170 Figure 2: IVD degeneration is quick and sustained following bilateral puncture of the caudal IVD.

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171 Representative Safranin-O stained sections of a representative control and the least and most degenerated

injured IVD from (A) sham and (B) bilateral puncture mice at 2, 4 and 12 weeks. Histopathological

grading of IVD degeneration ranges from moderate to severe with collapse of the IVD following

puncture. (C) Total degenerative grade was significantly increased at 2, 4 and 12 weeks after injury

compared to internal control which was not seen in sham. (D) Compartmentalized difference in
 degeneration between control and stab IVDs was seen in the nucleus pulposus, annulus fibrosus and at the

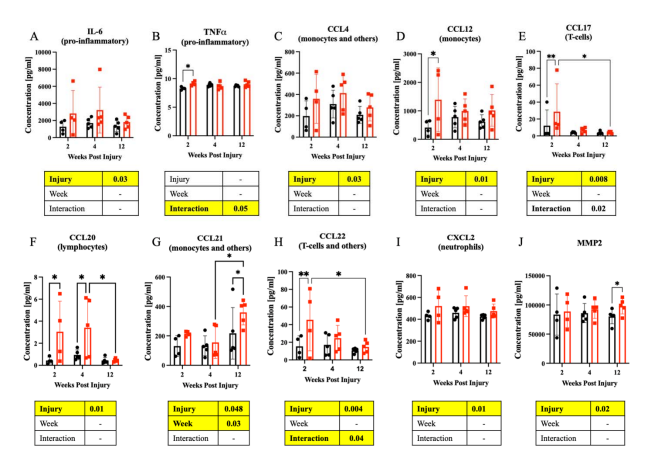
interfaces (a indicates p < 0.05 compared to internal controls). Data (A and B) were statistically analyzed

via paired two-way ANOVA with post-hoc Turkey HSD for control vs. injured. \* indicates p < 0.05. \*\*

indicates p < 0.01; \*\*\*\* indicates p < 0.001 in posthoc comparison. Scale bar is 500 µm.

- 180 3.2 Chemokine production peaks at 2 weeks after injury
- 181 Forty-four distinct chemokines and five MMPs were measured from the culture media of control and
- 182 punctured functional spinal units from injured mice. An effect of injury was seen in both pro-
- inflammatory chemokines (IL6 and TNFα) and immune cell recruitment chemokines (CCL4, CCL12,
- 184 CCL17, CCL20, CCL21, CCL22 and CXCL2) (Fig 3). TNFα and IL1β are pivotal inflammatory
- 185 chemokines in IVD degeneration, and in this experimental model we only see increased
- 186 TNF $\alpha$  expression.<sup>38,39</sup> The greatest difference between injured and control IVDs in chemokine production
- 187 occurred 2 weeks after injury where significantly higher expression of CCL12, CCL17, CCL20, CCL22
- and TNFa with all but CCL20 returning to control levels by 4 weeks post injury and CCL20 by 12 weeks
- 189 post injury (Fig 3A-E). CCL21 was elevated at 12 weeks after injury (Fig 3G). MMP-2 was detected as
- being affected by injury with the peak at 12 weeks post injury (Fig. 3J).





192

193 Figure 3: The injured IVD increased chemokine and MMP production. Chemokines measured can

broadly be characterized into 2 categories: pro-inflammatory and immune cell recruitment. Pro-

- inflammatory chemokines (A) IL-6 appears chronically elevated with injury while (B) TNF peaks at 2
- 196 weeks following injury. Immune cell recruitment chemokines are further categorized by their canonical
- 197 functions. There were also increases in chemokines associated with monocytes, (C) CCL4, (D) CCL12
- and (G) CCL21, with T-cells, (E) CCL17 and (H) CCL22 and lastly, (F) CCL20 which signals to
- 199 lymphocytes and (I) CXCL2 to neutrophils. (J) MMP-2 was the only of the five MMPs measured that was
- elevated with injury. \* indicates p < 0.05. \*\* indicates p < 0.01; \*\*\*\* indicates p < 0.001 in posthoc</li>
  comparison.
- 202 203
- 204 3.3 Innervation and vascularization propagate at different temporal trajectories
- 205 PGP9.5+ neurite and EMCN+ vasculature structures were manually segmented on a maximum projection
- 206 image (Fig 4A). The region of interest (ROI) contained the anterior and posterior outer annulus fibrosus
- and surrounding tissue. High magnification ROIs show innervation and vascularization that colocalize in
- these areas (Fig 4B). Nerve and vessel structures were semi-automatically traced and lengths were
- tabulated in each IVD.

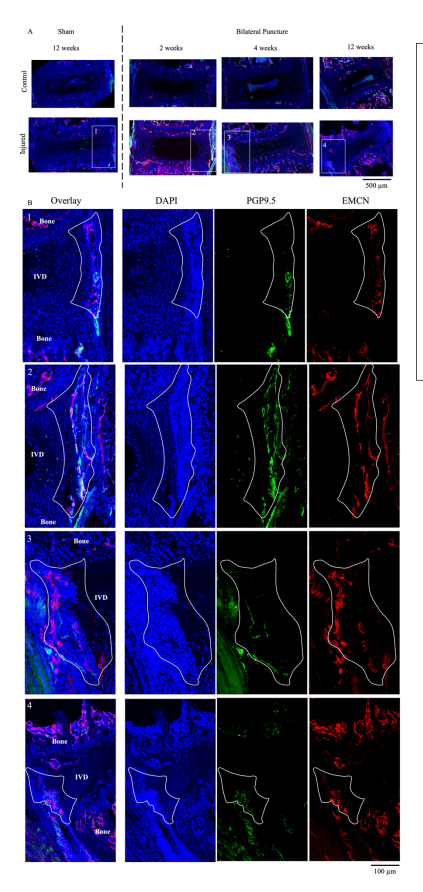
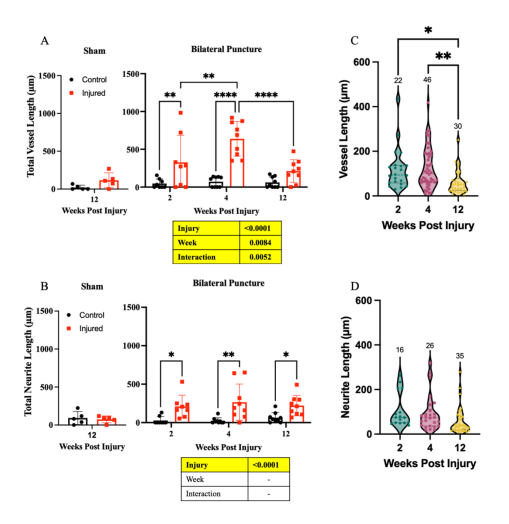


Figure 4: <u>Representative images</u> showing infiltration of neurites and vessels. (A) Maximum projection immunofluorescence images stained with PGP9.5 (Green) and EMCN (Red) against DAPI (blue) from sham and bilateral puncture mice. Control and injured are paired samples. Both the medial and lateral sides of the IVD are semi-automatically traced in ImageJ and used for further analysis. (B) Zoomed in regions from injured IVDs (sham and bilateral puncture). ROI, circled in white, includes the outer annulus fibrous (infiltrating and on the periphery) and surrounding disorganized tissue.

211 Sham mice showed negligible amounts of innervation and vascularization within their IVDs. Total length 212 of each feature in each IVD was measured and an increased presence of both structures was observed as early as 2 weeks after injury (Fig 5A-B). PGP9.5+ neurite structures are observed 2 weeks following 213 214 injury and remain consistently increased through the 12 week period; in comparison, EMCN+ vessels 215 peak at 4 weeks and appear to recede by 12 weeks after injury. Violin plots of punctured IVDs from injured mice show the tabulation of individual nerve and vessel lengths that were measured with the total 216 217 number of structures written above the plot (Fig 5C-D). The distribution of nerves remains consistent through all 12 weeks while the number of vessels in the 150-300 µm range is dramatically reduced at 12 218 219 weeks compared to 4 weeks post injury.



220

221 Figure 5: Quantification of feature lengths reveal divergent temporal progression of vessels and neurites.

All individual vessels and neurites were semi-automatically traced and their lengths tabulated. Total

223 length (sum of all individual lengths) of (A) vessels and (B) nerves demonstrate the presence of both

structures as early as 2 weeks after injury. Infiltrating vessels peak at 4 weeks and recede out to 12 weeks

after injury compared to nerves which remain consistently present through all 12 weeks. Violin plots of

the bilateral punctured IVDs of (C) vessel and (D) nerve length seems to demonstrate that the decreased total vessel length is mainly due to a reduced number of longer vessels compared to both 2 and 4 weeks

after injury. Number of vessels/nerves traced is indicated above each violin plot. \* indicates p < 0.05. \*\*

indicates p < 0.01; \*\*\*\* indicates p < 0.001 in posthoc comparison.

## 230 Discussion

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231 IVD injury models are a commonly utilized tool for studying the progression of IVD degeneration. In contrast to the lumbar spine, where degeneration is known to evoke changes in pain behavior<sup>24,40</sup>, we did 232 233 not observe any behavioral changes following injury in the caudal spine (Fig. S1). Thus, the caudal IVD 234 injury model is perhaps best suited to evaluate IVD-specific responses during degeneration. The advantage of a non-invasive surgical approach and access to multiple levels promotes the reduction in 235 number of research animals used in accordance to the 3R principle.<sup>41</sup> The use of the caudal spine also 236 minimizes the interference due to the disruption and inflammation of surrounding tissues compared to the 237 complex surgical access to the lumbar spine. While there have been extensive studies showing the 238 structural and compositional degenerative changes following caudal puncture<sup>25,29,30,42,43</sup>, limited data 239 exists on additional aspects of IVD degeneration including innervation<sup>27,28</sup> and vascularization and 240 241 chemokine secretion from the explanted IVD.<sup>44</sup> Bilateral puncture of the caudal IVD resulted in quick and sustained IVD degeneration up to 12 242 weeks post-injury. Both proinflammatory (IL6 and TNFa) and chemokines (CCL4, CCL12, CCL17, 243 CCL20, CCL21, CCL22 and CXCL2) were elevated with injury, with the highest expression of a subset 244

of chemokines compared to controls at 2 weeks following injury. These chemokines canonically recruit
 monocytes, T-cells, and lymphocytes (Fig.3).<sup>45</sup> Yet chemokines are known to be pleiotropic and have
 been associated with additional functions such as IVD degeneration, pain, neurite growth and

angiogenesis.<sup>46</sup> For example, CCL4 has been shown to be elevated in degenerated human IVDs and

250 lumbar IVDs has been shown to enhance pain behavior changes, possibly through irritation of nerve

associated with pain behavioral changes in a rat model of IVD degeneration.<sup>6,47,48</sup> TNF injected into rat

endings.<sup>33</sup> CCL17 and CCL21 induced dorsal root ganglion (DRG) axonal growth <sup>49,50</sup> and CXCL2 is a

known mediator of angiogenesis.<sup>51</sup> CCL17 and CCL22 through the receptor CCR4 were indicated to play

a role in pain development and CCL22 was able to activate neurons and increase neuron excitability.<sup>52</sup>

254 Chemokines production of the IVD following injury may help provide further insights into the

pathoanatomy of innervation and vascularization as well as provide possible pathways for IVD
 degeneration associated low back pain.

Innervation of the IVD may be the potentiator of low back pain observed with lumbar puncture 257 258 models and this feature is recapitulated here in the caudal spine. Studies have previously illustrated 259 innervation of the IVD following injury with detection of PGP9.5+ or CGRP+ staining injured IVDs, but without any quantification of the structures.<sup>19,26,28</sup> Further, the coincidence of vascularization with neo-260 innervation has been previously observed but the time course of vessel propagation into the IVD 261 following injury has not been documented. To address these previous limitations, we semi-automatically 262 traced neurite and vascular structures on maximum projection images of PGP9.5 and EMCN stained thick 263 sections.<sup>37</sup> This allowed for the tabulation of neurites and vessels present in the region of interest and their 264 265 lengths for comparison where we saw a time-dependent vessel but not neurite infiltration of the outer 266 annulus fibrosus and surrounding tissue. Penetration of the IVD by vessels would be considered prerequisite to infiltration by circulating cells, including monocytes and other immune cells that might be 267 responsible for secretion of the chemokines. 268

Behavioral assays can be used following lumbar puncture to quantify pain.<sup>20,23,24,26</sup> A limitation of 269 270 the caudal puncture model is that we do not expect to see behavioral changes that would be traditionally 271 associated with low back pain. No differences in behavioral measures between sham and bilateral 272 puncture mice were detected (Fig. S1). Alternative measures of pain can include sensitization of the tail to mechanical and thermal stimuli (e.g., Hargreave's test, tail-flick).<sup>28,53</sup> Another possible surrogate of 273 274 pain-related change is to quantify molecular expression of neurotrophic factors in the innervating lumbar dorsal root ganglia (DRG). The DRG has been linked to chronic pain, and the increase in the expression 275 of pain-related neuropeptides as well as neuronal excitability may be the mediators of discogenic pain.<sup>54</sup> 276 277 Ongoing work utilizes immunohistochemical staining for altered presence of neurotrophic factors in 278 lumbar DRGs that have a demonstrated role in mediating pain transmission in the spine such as transient 279 receptor potential cation channel subfamily V member 1 (TRPV1) (Fig. S2).

In this study we used utilized a more sophisticated method to quantify innervation and 280 vascularization of the IVD than has been previously done in puncture models. Protein analysis the IVD 281 secreted chemokines revealed potential molecular mediators of IVD degeneration, innervation and 282 283 vascularization with relevance to generation of inflammation and pain. Many of the secreted chemokines 284 found to be elevated may be associated with increased presence of infiltrating monocytes that may include macrophages, B-cells or T-cells.<sup>55,56</sup> Not surprisingly, the key angiogenic factor, VEGFA, was not 285 286 elevated at any of the measured timepoints. VEGFA is critically expressed early following tissue repair promote early angiogenesis, <sup>55</sup> and by two-weeks following injury VEGFA has already exerted its effects 287 as evidenced by robust vessel formation. In this study, we are limited to chemokines differentially 288 expressed between the injured and control IVDs; thereby, excluding chemokines that are more 289 systemically expressed following puncture. Recent work shows that the chronic NF $\kappa$ B activation in the 290 caudal IVD produces a secretome that promote macrophage migration.<sup>57</sup> Our data here confirm that a 291 degeneration-causing injury will upregulate a plethora of chemokines that will likely recruit multiple 292 immune cell types<sup>55</sup>, concomitant with increasing neurovascular features. Future studies quantifying the 293 294 presence of these immune cells would advance our understanding of a role for local versus systemic changes in modulating chemokine secretion, as well as key factors that govern the infiltration of these 295 296 pain-associated features.

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300

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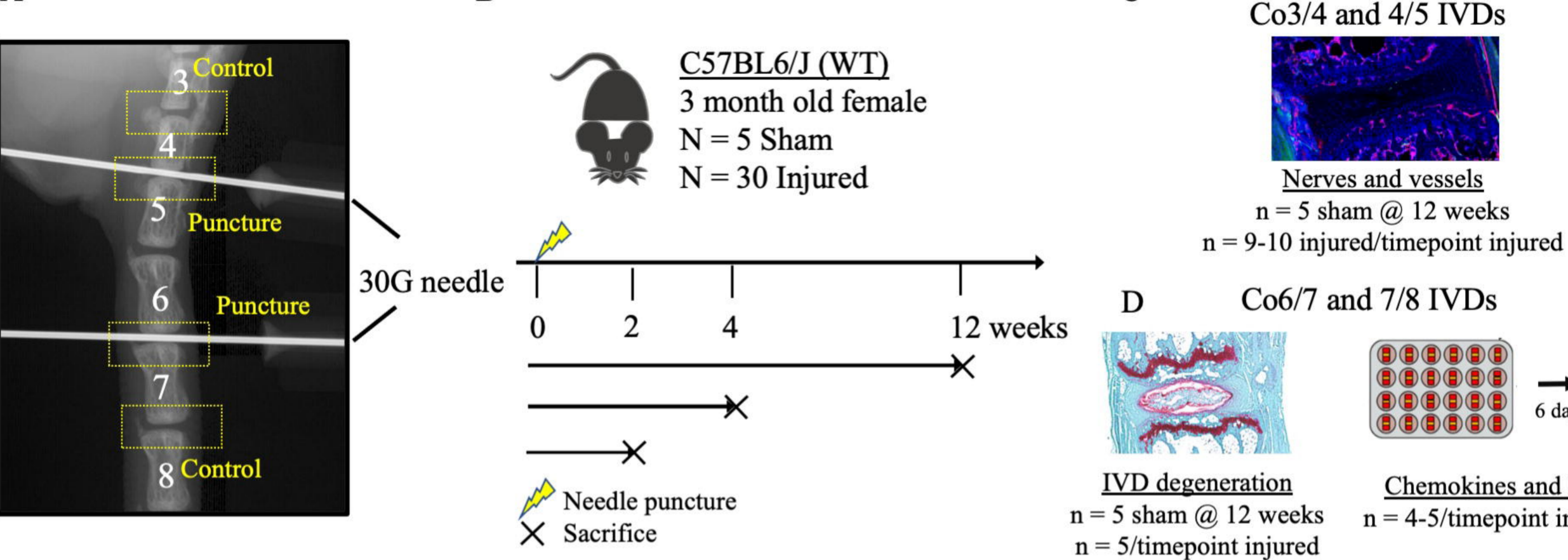
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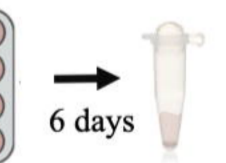


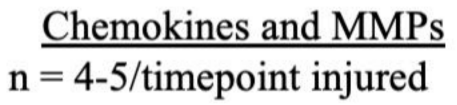


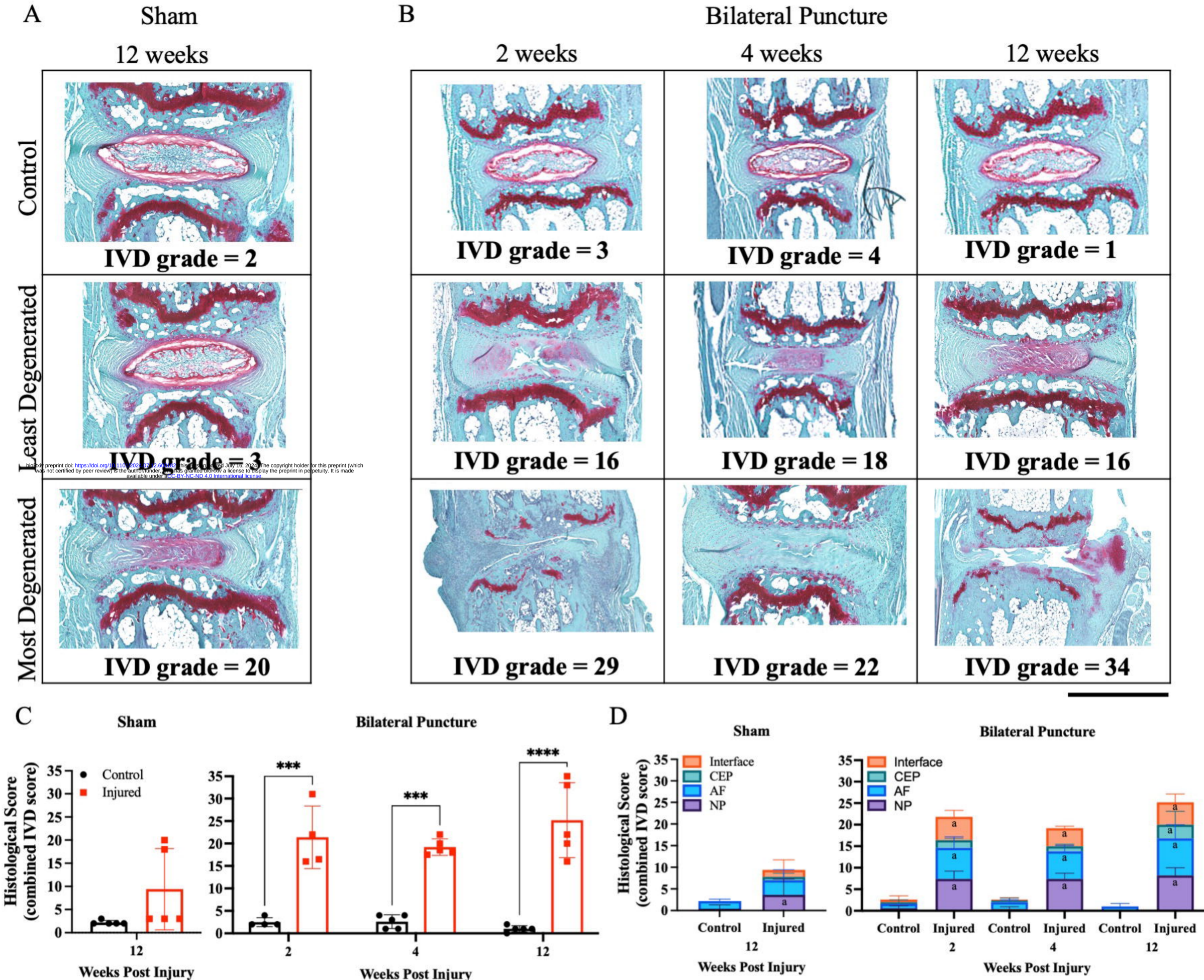
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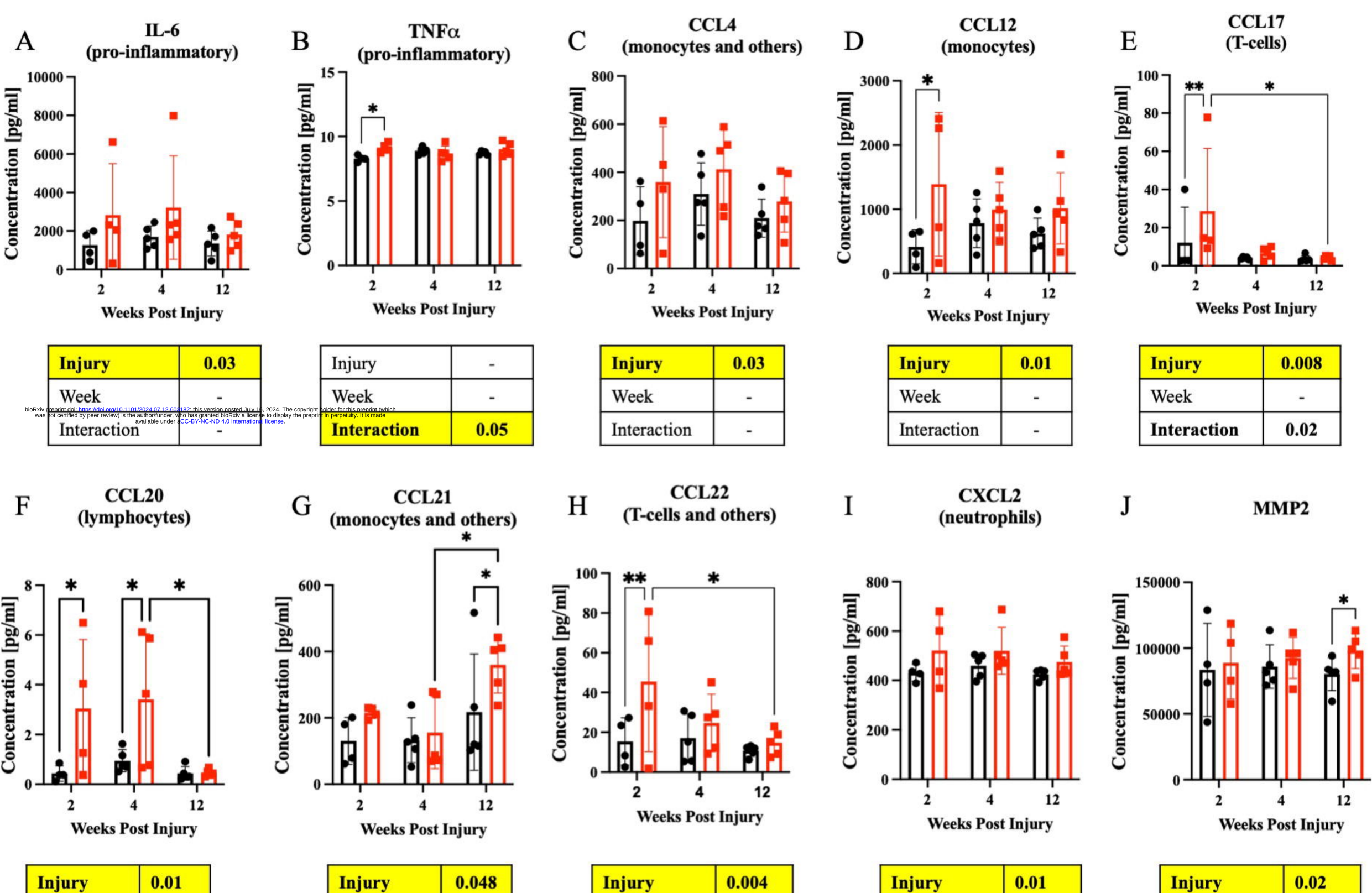


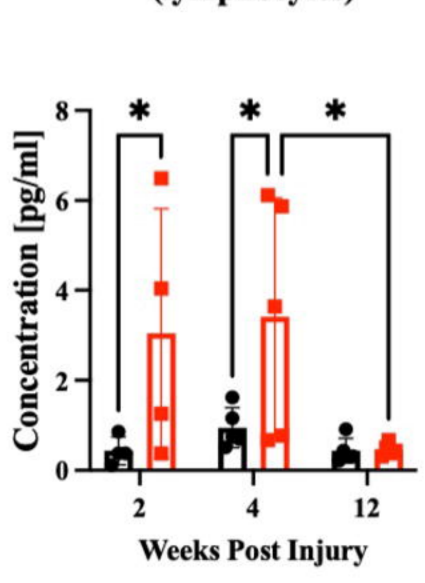




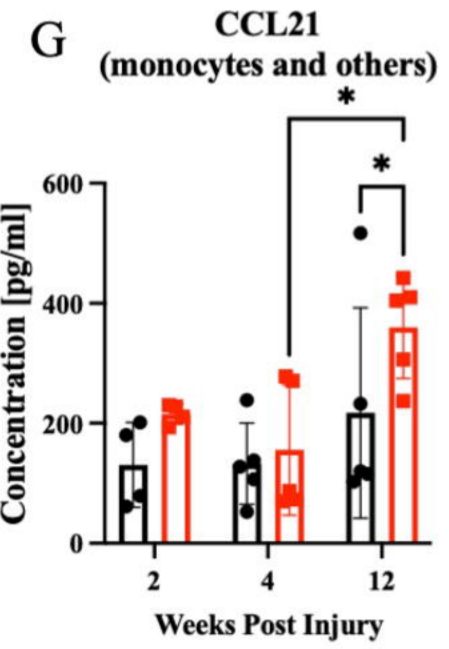


Weeks Post Injury

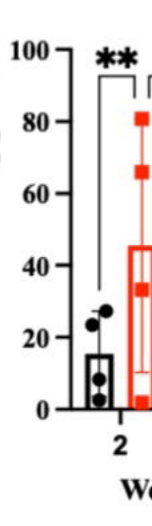




Injury	0.01
Week	-
Interaction	6 <del></del> 0



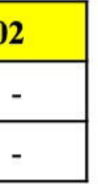
Injury	0.048
Week	0.03
Interaction	-

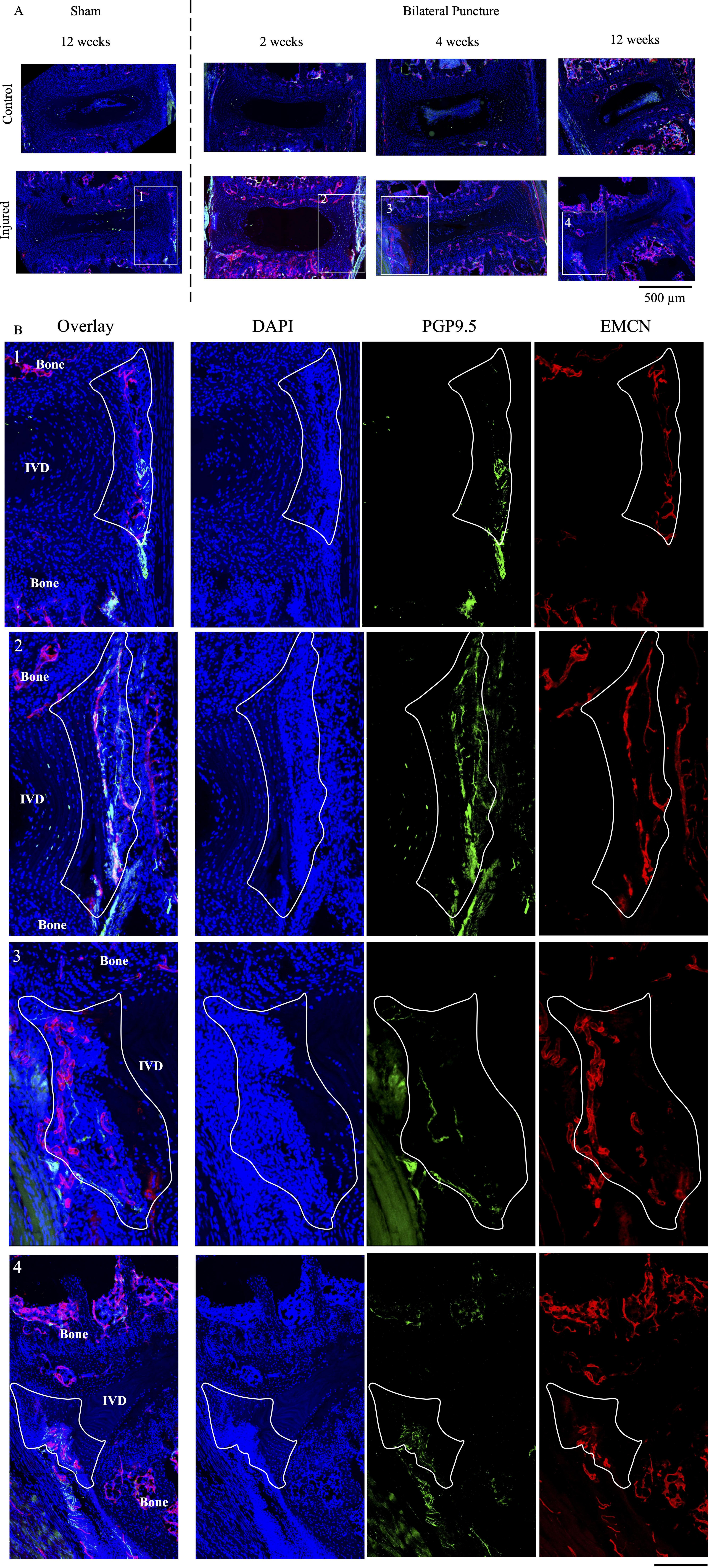


Injury	0.004
Week	-
Interaction	0.04

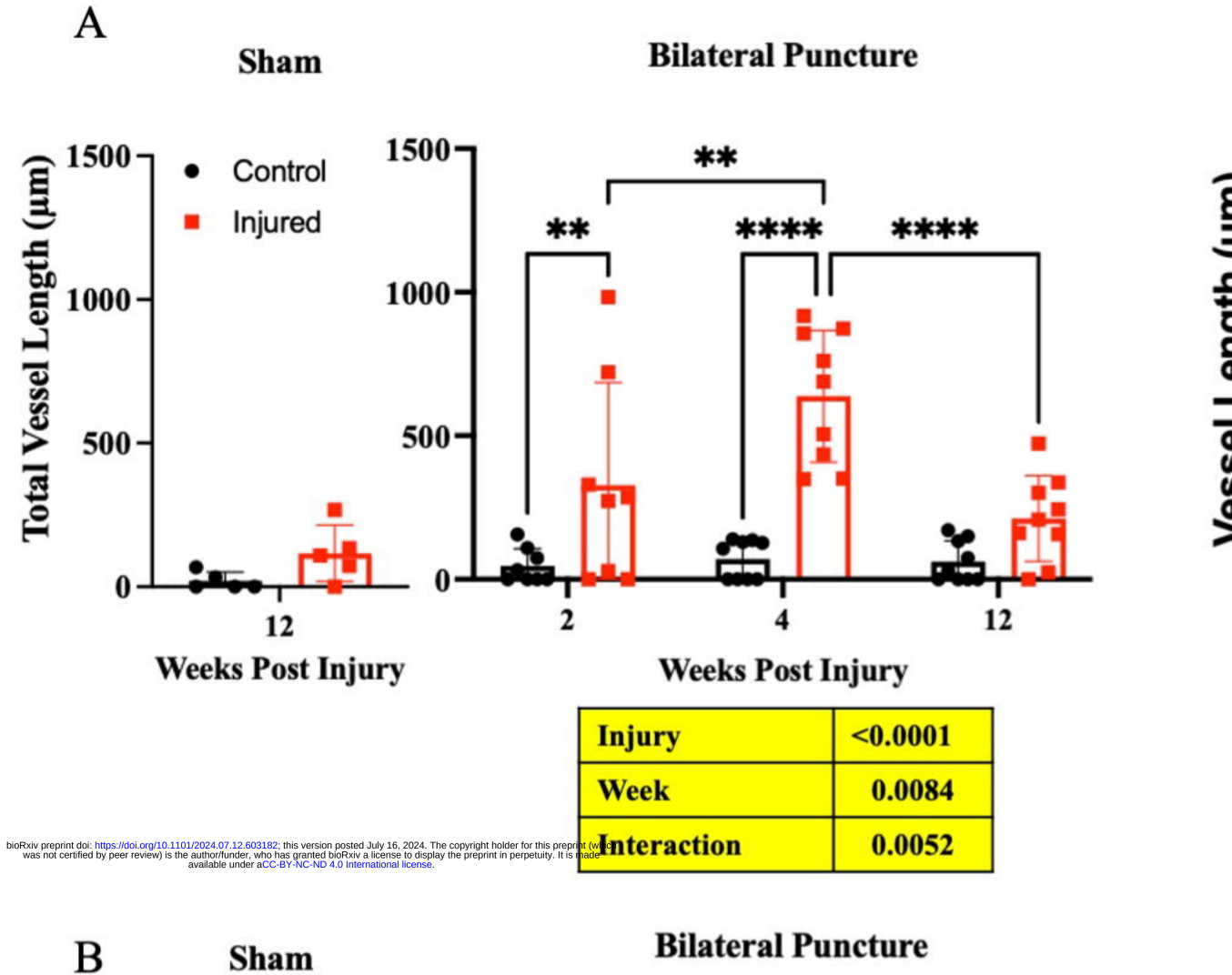
Week -Interaction -

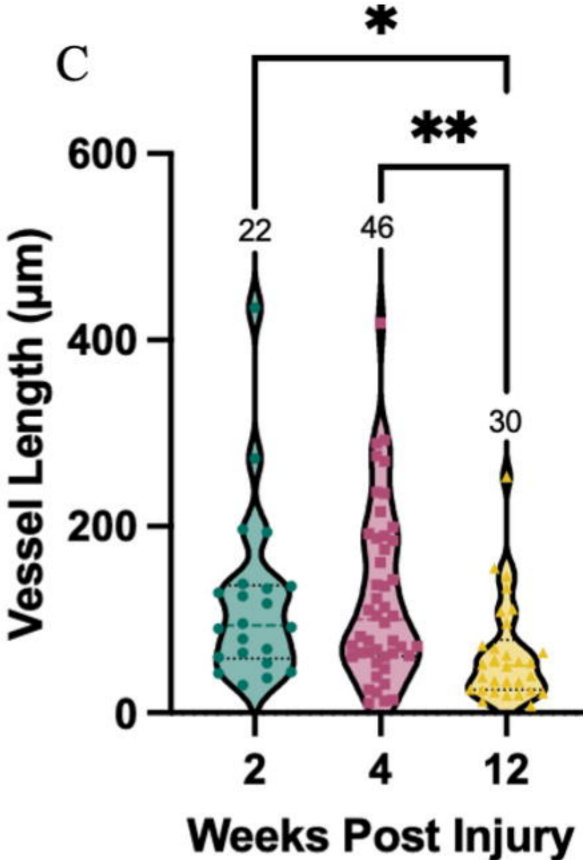
Injury	0.0
Week	
Interaction	





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