1 Microglial refinement of A-fibre projections in the postnatal spinal cord dorsal

#### 2 horn is required for normal maturation of dynamic touch.

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#### 18 Impact statement

- 19 Microglia phagocytose superfluous A-fibres in the superficial spinal dorsal horn
- 20 during normal development, the disruption of which leads to long term aberrant
- 21 dynamic touch processing and behaviour.

#### 22 Abstract

23 Sensory systems are shaped in postnatal life by the refinement of synaptic 24 connections. In the dorsal horn of the spinal cord, sensory circuits undergo postnatal 25 activity dependent reorganisation, including the retraction of primary afferent A-fibres 26 from superficial to deeper laminae which is accompanied by decreases in cutaneous 27 sensitivity. Here we show that microglia, the resident immune cells in the CNS, 28 phagocytose A-fibre terminals in superficial laminae in the first weeks of life. Genetic 29 perturbation of microglial engulfment at that time prevents the normal process of A-30 fibre retraction, resulting in increased sensitivity of dorsal horn cells to dynamic 31 tactile cutaneous stimulation, and behavioural hypersensitivity to dynamic touch. 32 Thus, functional microglia are necessary for normal postnatal development of dorsal 33 horn sensory circuits. In the absence of microglial engulfment, superfluous A-fibre 34 projections remain in the dorsal horn and the balance of sensory connectivity is 35 disrupted, leading to lifelong hypersensitivity to dynamic touch.

36

#### 38 Introduction

39 The neonatal spinal dorsal horn differs substantially from that in adults and 40 undergoes extensive structural and functional reorganisation over the postnatal 41 period. One notable change is the termination zone of primary afferent A-fibres, the 42 large myelinated afferents that encompass many low threshold cutaneous tactile 43 afferents<sup>1</sup>. These afferent terminals occupy both superficial and deep laminae of the 44 dorsal horn in neonatal rodents and gradually retract to terminate in deeper laminae III-IV by the end of the 3rd postnatal week and in adulthood  $^{2-4}$ . This retraction of A-45 46 fibre terminals is accompanied by a reduction in dorsal horn cell receptive field sizes 47 on the skin, as well as a decline in tactile sensitivity and an increase in reflex 48 behaviour precision, with similar changes in somatosensory behaviour observed in 49 human infants (Fitzgerald 1985, 2015; Fitzgerald et al. 1988). This suggests that 50 structural refinement in the dorsal horn likely underlies behavioural maturation. 51 The process of A-fibre terminal retraction in the first weeks of life is activity 52 dependent: Blocking neuronal input through spinal NMDAR inhibitors or increasing 53 the noise of neuronal input through random vibration to the skin over extended period both prevented the normal retraction of A-fibres<sup>8,9</sup>. However, the exact 54 55 mechanism underlying the retraction of A-fibre terminals is not known. Microglia 56 cells, the major phagocytes in the CNS, have been shown to remove superfluous 57 neurons both by driving apoptosis, removing apoptotic cells, and phagocytosing synapses and neurites during postnatal refinement <sup>10</sup>. To date, such studies have 58 been largely restricted to the brain <sup>11–15</sup>, with two studies reporting a role of microglia 59 in the postnatal development of spinal cord ventral horn motor circuits <sup>16,17</sup>. Whether 60 61 microglia are also involved in the maturation of somatosensory circuits in the dorsal 62 horn is not known. Here we hypothesise that the retraction of A-fibres from

superficial laminae in the postnatal period is driven by microglia which prune A-fibre
terminals in the dorsal horn as part of normal postnatal development of spinal
sensory circuits.

66 Microglia undergo postnatal maturation during which they not only change in density and morphology but also alter their transcriptional and functional identity <sup>18,19</sup>. Brain 67 68 microglia show particularly high expression of lysosome associated genes at P4/5 suggesting a specialised role of microglial phagocytosis during development <sup>20</sup>. In 69 70 addition, microglia exhibit spatial heterogeneity, as they populate different brain 71 regions at different rates postnatally, and express distinct local genetic profiles and 72 phenotype in adulthood <sup>21-23</sup>. So far, most research has focused on the brain and 73 relatively little is known about spinal cord microglia.

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75 To test whether A-fibre terminals in the developing spinal dorsal horn are pruned by 76 microglial phagocytosis, we used a transgenic mouse line that expresses tdTomato (tdT) under the Valut1 promoter which labels a subset of A-fibres <sup>24</sup> and mapped 77 morphological changes in dorsal horn microglia across the first two postnatal weeks 78 79 using immunofluorescence and confocal microscopy. We next tested whether normal 80 microglial phagocytosis was required for the A-fibre pruning. Constitutive knock-out 81 of the gene *Tmem16f* in microglia was shown to reduce microglial phagocytosis and 82 motility in the adult spinal cord in a neuropathic pain model, resulting in reduced pain 83 behaviour<sup>25</sup>. Therefore, we blocked microglial phagocytosis during the first postnatal 84 week using a tamoxifen inducible Cre-mediated deletion of the Tmem16f gene in 85 microglia and measured the effects on A-fibre pruning, the subsequent maturation of 86 dorsal horn synaptic connections, and behavioural reflex sensitivity to mechanical 87 skin stimulation. The results suggest that microglia prune A-fibre terminals in the

- 88 developing spinal dorsal horn and that this postnatal microglial refinement of A-fibre
- 89 terminals is required for normal somatosensory maturation.
- 90

91 **Results** 

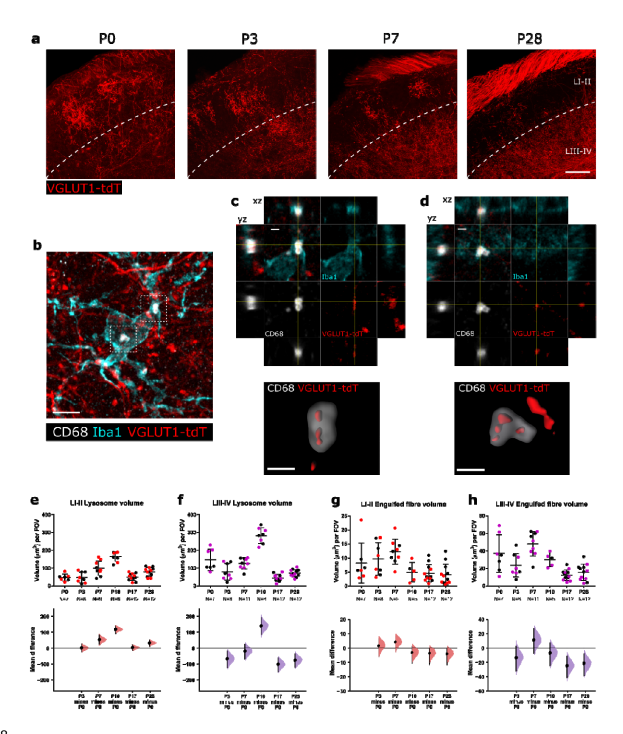
# 92 Dorsal horn microglia phagocytose A-fibre terminals during postnatal 93 development

94 In the developing spinal cord, tactile-encoding A-fibres initially project throughout the 95 dorsoventral extent of the dorsal horn and refine over the first few postnatal weeks to 96 terminate in deeper laminae, segregated from the more superficial terminals of 97 noxious-encoding C-fibres<sup>8</sup>. We confirmed this using a transgenic reporter mouse in 98 which tdTomato is expressed in vesicular glutamate transporter 1 (Vglut1) 99 expressing neurons (VGLUT1-tdT), a subpopulation of large myelinated sensory 100 neurons with features consistent with Aβ-low threshold mechanoreceptors (LTMRs) <sup>24</sup>. This tdT-expression was sparse at P0 and increased with age, likely due to the 101 increasing developmental expression profile of Vglut1 26,27 (Figure 1-figure 102 103 supplement 1a). Despite this, the mice clearly revealed A-fibre terminals with flame-104 shaped arbors extending into laminae I-II up to P7 (Figure 1a), which are no longer present in these superficial laminae by P28, consistent with previous findings<sup>8</sup>. 105

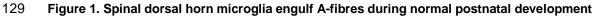
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We hypothesised that microglia might be involved in this refinement process by removing A-fibres in the superficial laminae via phagocytosis and determined the phagocytic activity of dorsal horn microglia over this period using the lysosome associated molecular marker CD68 (Figure 1b-f, Figure 1—figure supplement 1i-k). CD68 lysosome volume increased in both laminae I-II and III-IV over the postnatal

112 period, reaching a peak at P10 and declining thereafter at P17 and P28 (LI-II P10 vs 113 P0 unpaired mean difference 117.16 [95.00% CI 92.26, 137.61], LIII-LIV unpaired 114 mean difference 136.02 [95.00% CI 82.89, 184.41]. It is known that the dorsal horn 115 undergoes developmental apoptosis around birth, extending into the first postnatal 116 week. Therefore, to determine whether the rise in CD68 could be contributed to 117 engulfment of apoptotic cells, we also quantified the number of apoptotic cells and 118 phagocytic cups per microglia. In contrast to the rise in CD68, the number of 119 phagocytic cups per microglia decreased over the first postnatal week concomitant 120 with a decrease in apoptotic cell counts (Figure 1—figure supplement 2), suggesting 121 that the increase in CD68 is not due to phagocytosis of apoptotic cells but other 122 material (Phagocytic cups: F(3, 12)=10.97, P=0.0009, apoptotic cells: F(3, 123 12)=14.18, P=0.0003). In parallel, microglial density and their ratio to neurons 124 increased over this period (Figure 1-figure supplement 1b-h). Together this 125 suggests that the first postnatal week represents a distinct period of microglial 126 activity, characterised by high levels of microglial CD68.







a. VGLUT1-tdTomato (red) expression in spinal laminae I-II decreases across age. Dashed white line
 indicates border between lamina II and lamina III. Scale bar = 50µm.

**b.** Representative z-projected super-resolution image of A-fibre engulfment by microglia within the cell body. White inset box show location of higher magnification panels in **c** and **d**. Scale bar = 5  $\mu$ m.

c, d. High magnification images of microglial A-fibre engulfment in b stained for microglia (Iba1, cyan),
 microglial lysosomes (CD68, white) and endogenously fluorescent A-fibres (VGLUT1-tdT, red). Cross hairs show position of the xz and yz side-view panels. Bottom panels show surface rendering of the
 super-resolution image revealing pieces of tdT labelled fibres engulfed inside lysosome. Scale bar = 1

138 µm.

**e.** Microglial lysosome volume peaks at P10 and decreases thereafter for LI-II. P3 vs. P0: mean difference 0.48 [95% CI 23.52, 24.35]; P7 vs. P0: mean difference 52.19 [95% CI 23.71, 79.47]; P10 vs. P0: mean difference 117.16 [95% CI 92.26, 137.61]; P17 vs. P0: mean difference 2.52 [95% CI - 15.07, 18.38]; P28 vs. P0: mean difference 31.84 [95% CI 11.42, 49.16]. Field of view (FOV) = 245 $\mu$ m x 65 $\mu$ m.

144**f.** Microglial lysosome volume peaks at P10 and decreases thereafter for LIII-IV. P3 vs. P0: mean145difference -66.67 [95% CI -121.07, -18.14]; P7 vs. P0: mean difference -19.48 [95% CI -67.54, 21.66];146P10 vs. P0: mean difference 136.02 [95% CI 82.89, 184.41]; P17 vs. P0: mean difference -100.85147[95% CI -145.40, -62.42]; P28 vs. P0: mean difference -74.95 [95% CI -119.97, -36.61]. Field of view148(FOV) = 245 $\mu$ m x 65 $\mu$ m.

149 g. Engulfed fibre volume peaks at P7 and decreases thereafter for LI-II. P3 vs. P0: mean difference
150 1.46 [95% CI -6.86, 6.29]; P7 vs. P0: mean difference 4.05 [95% CI -4.11, 8.18]; P10 vs. P0: mean
151 difference -3.28 [95% CI -10.76, 1.11]; P17 vs. P0: mean difference -3.69 [95% CI -11.57, -0.10]; P28
152 vs. P0: mean difference -4.11 [95% CI -11.79, -0.38]. Field of view (FOV) = 245µm x 65µm.

h. Engulfed fibre volume peaks at P7 and decreases thereafter for LIII-IV. P3 vs. P0: mean difference
-13.47 [95% CI -31.97, 2.15]; P7 vs. P0: mean difference 11.03 [95% CI -7.37, 26.33]; P10 vs. P0:
mean difference -7.05 [95% CI -24.52, 6.62]; P17 vs. P0: mean difference -24.55 [95% CI -40.82, 11.27]; P28 vs. P0: mean difference -21.49 [95% CI -37.93, -7.92]. Field of view (FOV) = 245µm x
65µm.

158 N-numbers as indicated. Black and colourful data points indicate females and males respectively.

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160 We next asked whether the high level of CD68 expression in microglia during the 161 first postnatal week is associated with the refinement of neural connectivity in the 162 dorsal horn through the removal of superfluous A-fibre terminals. Engulfment of A-163 fibres was measured by quantifying VGLUT1-tdT fluorescence volume within CD68-164 positive lysosomes inside lba1-labelled microglia cells (Figure. 1 b-d, g-h). 165 Consistent with the postnatal increase in microglial lysosome volume, the volume of 166 engulfed A-fibres was high during the first postnatal week, peaking at P7 and 167 decreasing thereafter in both laminae I-II, (unpaired mean difference P28 vs. P0 -168 4.11 [95% CI -11.79, -0.38]) and laminae III-IV (unpaired mean difference P28 vs. P0 169 -21.49 [95% CI -37.93, -7.92]). This effect was more pronounced in the deeper 170 laminae LIII-IV, which has higher volumes of fibre engulfment at all ages compared 171 to laminae I-II likely due to the higher density of A-fibres in the deeper laminae.

#### 172 Microglial activity is required for normal A-fibre pruning in laminae I-III

173 Having established that microglia participate in postnatal dorsal horn remodeling of 174 A-fibre afferent terminals, we sought to determine whether microglial function is 175 necessary for this process to occur normally. The phospholipid scramblase TMEM16F is required for normal microglial phagocytic activity<sup>10</sup>. To address this 176 177 question we disrupted microglial function postnatally using a Cre-inducible conditional knocking out of Tmem16f in microglia cells <sup>25</sup>, and observed the 178 structural, functional and behavioural consequences in adult animals. A R26<sup>LSL-Ai9</sup> 179 reporter line was used to label Cxc3cr1<sup>Cre</sup>-expressing microalial cells <sup>28,29</sup> and 180 *Thy*  $1^{eGFP}$  allele was used to identify A-fibres <sup>30,31</sup>. 181

To control for off-target effects of Cre expression and tamoxifen administration, both cKO mice and control mice ( $Cx3cr1^{CreER/+}$ ;  $Tmem16f^{I/fl}$ ;  $R26^{LSL-Ai9}$ ;  $Thy1^{eGFP}$  and  $Cx3cr1^{CreER/+}$ ;  $Tmem16f^{+/+}$ ;  $R26^{LSL-Ai9}$ ;  $Thy1^{eGFP}$  respectively) received 4hydroxytamoxifen (4-HT) daily from P1 to P3 and were assessed at 3-4 months old. Specificity of Cre expression was confirmed with  $R26^{LSL-Ai9}$  tdT expression.

# Adult microglial Tmem16f cKO mice have increased VGLUT1+ terminals in superficial but not deep laminae

Postnatal deletion of microglial *Tmem16f* resulted in adults with increased A-fibre terminal occupancy in the superficial dorsal horn, as revealed by both Thy1-GFP expression and VGLUT1 immunohistochemistry (Figure 2a-b, d). As Thy1-GFP is expressed in only a small number of sensory neurons and varied across animals, all subsequent quantification was performed using VGLUT1 immunolabelling. Synaptic density of this increased input was quantified using VGLUT1 immunohistochemistry in laminae I-III (Figure 2a upper panels). *Tmem16f* cKO mice had greater primary

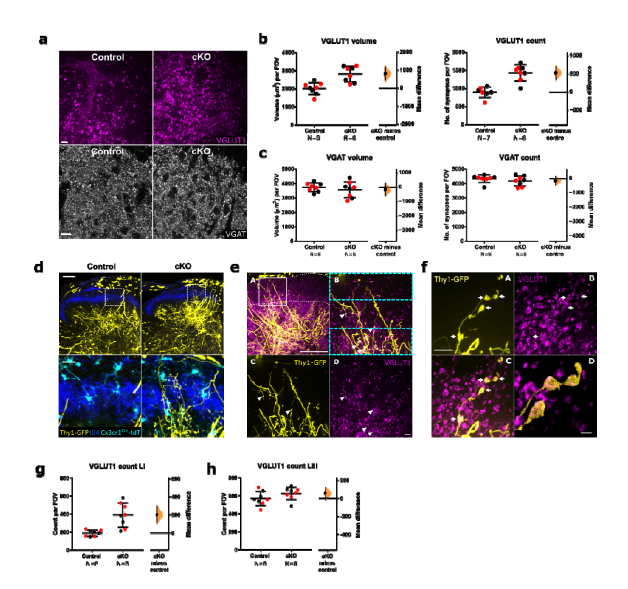
afferent VGLUT1 synaptic density throughout the dorsal horn than control mice as
measured by both total synapse volume and synapse number (Figure 2b) (unpaired
mean difference VGLUT1 volume: 799.28 [95.00% CI 450.09, 1136.35], VGLUT1
number: 533.67 [95.00% CI 356.12, 695.50]). In contrast, local inhibitory VGAT
synapse density was unaltered (Figure. 2a, c) (unpaired mean difference VGAT
volume: -168.31 [95.00% CI -562.61, 253.44], VGAT number: -159.19 [95.00% CI 416.42, 148.88]).

As synaptic engulfment was also observed in the deeper laminae we re-analysed the data by cropping the original images to only contain LI or LIII to assess potential differences across laminae (Figure 2e). Surprisingly, VGLUT1 density in LIII was unaltered in contrast to LI (Figure 2g-h), suggesting that the changes observed are mainly driven by an increase in superficial VGLUT1 terminal projections (unpaired mean difference superficial/LI VGLUT1 count: 201.33 [95.00% CI 110.90, 292.00], deep/LIII VGLUT1 count: 56.61 [95.00% CI -13.81, 118.42]).

To investigate whether the superfluous VGLUT1 terminals indeed form synaptic contacts, we also co-stained spinal cord sections from *Tmem16f* cKO and control animals with HOMER, which marks postsynaptic densities. Indeed, we found colocalization of VGLUT1-positive Thy1-GFP labelled A-fibre terminals with HOMER were increased in *Tmem16f* cKO animals, suggesting that a surplus of functional VGLUT1 positive synaptic contacts are formed in the superficial dorsal horn of *Tmem16f* cKO animals (Figure 2—figure supplement 1).

Thus, targeted deletion of microglial *Tmem16f*, and consequently impaired phagocytosis specifically disrupted the normal developmental pruning of excitatory afferent A-fibre projections in the superficial dorsal horn.

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#### 222 Figure 2. Neonatal *Tmem16f* cKO in microglia increases dorsal horn A-fibre terminals.

a. Representative images of VGLUT1 and VGAT puncta from the spinal dorsal horn of adult *Tmem16f* control and cKO animals. Field of view (FOV) = 94µm x 94µm (VGLUT1), 96µm x 96µm (VGAT).
 Scale bars = 10µm.

226 **b.** VGLUT1 puncta volume and count were both increased in adult *Tmem16f* cKO animals compared

to controls. Control vs cKO volume: mean difference 799.28 [95.00% CI 450.09, 1136.35]; control vs
 cKO count: mean difference 533.67 [95.00% CI 356.12, 695.50].

229 c. VGAT puncta volume and count were not significantly different between adult Tmem16f cKO

animals and controls. Control vs cKO volume: mean difference -168.3105 [95.00% CI -562.61,

231 253.44]; control vs cKO count: mean difference -159.19 [95.00% CI -416.42, 148.88].

232 d. Thy1-GFP labelled A-fibres (yellow) are present in the superficial dorsal horn (delimited by IB4,

233 blue) of adult cKO mice, but not in adult controls. tdTomato-labelled microglia (cyan) are present in

both cKO and control animals. Lower panels show high power images of the boxed areas in the

235 upper panels. Scale bar =  $100\mu$ m in the top panel and  $20\mu$ m in the lower panel.

e. Thy1-GFP labelled A-fibres (yellow) in the superficial dorsal horn laminae of adult *Tmem16f* cKO
 mice express VGLUT1 (magenta). (A) Low magnification image showing superficially-projecting Thy1-

238 GFP labelled fibres. White box indicates where images were taken for analysis in **a-c**. (B) High

239 magnification of boxed region in A. Cyan boxes indicate the cropped areas used for analysis in g and

h. (C) Thy1-GFP labelling. (D) VGLUT1 immunoreactivity. Scale bar in A = 100μm. Scale bar in D =
 10μm.

242 f. A-D. High-magnification examples for VGLUT1-expression (magenta) in Thy1-GFP labelled A-fibres

243 (yellow) in adult *Tmem16f* cKO mice. Scale bar in A =  $20\mu$ m, scale bar in D =  $5\mu$ m.

g, h. VGLUT1 count in adult *Tmem16f* cKO and control animals in superficial lamina I (g) and deep

- 245 Iamina III (h). LI VGLUT1 count mean difference 201.33 [95.00% CI 110.90, 292.00]; LIII VGLUT1
- 246 count mean difference 56.61 [95.00% CI -13.81, 118.42]. FOV = 31µm x 94µm.

247 N-numbers as indicated. Black and red data points indicate females and males respectively.

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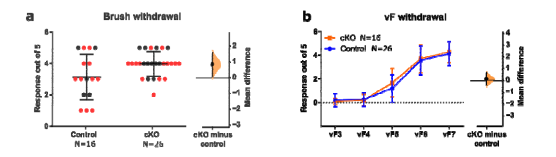
249 Dorsal horn sensory neurons in adult microglial Tmem16f cKO mice are less 250 responsive to dynamic tactile stimulation of the skin, but have larger receptive 251 fields

252 We reasoned that the reduced A-fibre pruning due to impaired neonatal microglial phagocytosis in Tmem16f cKO mice would alter behavioural sensitivity to hindpaw 253 254 tactile stimulation. The hindlimb withdrawal reflex was measured in response to 255 brushing of the plantar surface of the paw. Tmem16f cKO mice showed a greater 256 number of withdrawals in response to dynamic brushing than controls (unpaired mean difference 0.83 [95% CI 0.072, 1.57]), but not to static vF stimulation (Figure 3, 257 Figure 3—figure supplement 1). Tmem16f cKO animals therefore displayed 258 259 cutaneous hypersensitivity specifically towards dynamic low-threshold tactile 260 stimulation.

261 We hypothesised that excessive A-fibre terminals in dorsal horn might lead to 262 increased dorsal horn activity which could underlie the behavioural hypersensitivity. 263 To test this, we used *in vivo* single unit extracellular recordings in the dorsal horn of 264 anaesthetised Tmem16f cKO and control mice. We recorded from adapting and non-265 adapting wide dynamic range neurons (WDR) in the deep dorsal horn, which according to the criteria by Lee et al. 2019<sup>32</sup> are presumed to be excitatory and 266 267 inhibitory neurons respectively. Of 30 cells recorded from control mice, 8 were non-268 adapting and 22 were adapting. Of 28 cells recorded from Tmem16f cKO mice, 5

were non-adapting and 23 were adapting. This is consistent with the expected ratio of 1:2 of inhibitory to excitatory cells as reported in previous studies <sup>32–34</sup> (P=0.56 for control and P=0.11 for cKO in Binomial test against expected ratio). We quantified spinal neuronal response to 1) dynamic innocuous tactile stimuli (brush), and 2) static von Frey hair (vF) stimuli applied to the plantar hindpaw as well as mapped dynamic touch-sensitive receptive field areas on the plantar hindpaw.

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Figure 3. Neonatal *Tmem16f* deletion in microglia increases dynamic brush sensitivity
 a. Brush withdrawal response for *Tmem16f* cKO animals are higher than control animals, mean
 difference 0.83 [95% CI 0.072, 1.57].

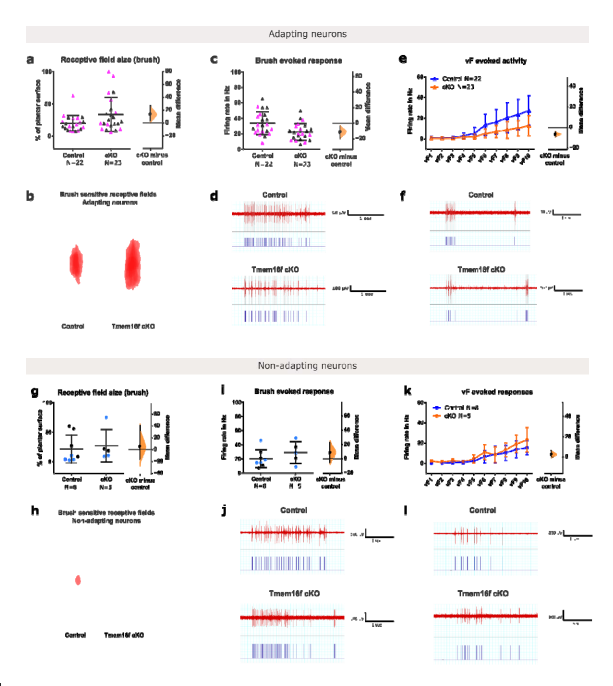
b. vF withdrawal response did not differ between *Tmem16f* cKO and control animals, mean difference
 0.12 [95% CI -0.39, 0.65].

- 282 N-numbers as indicated. Black and red data points indicate females and males respectively.
- 283

284 We reasoned that excessive A-fibre presence in the dorsal horn would increase their 285 spatial connectivity with postsynaptic neurons, leading to larger dorsal horn receptive 286 field areas. Consistent with increased A-fibre synaptic contacts, mean brush 287 receptive field sizes for adapting neurons were increased by almost 50% (unpaired 288 mean difference 13.7 [95.00% CI 3.3, 27.2]), with a notable subpopulation of cKO 289 neurons expanding their receptive fields to cover the entire, or majority of the plantar 290 surface (Figure 4a, b). The expansion of receptive field area was accompanied by a 291 reduced number of spikes in response to both brush and vF stimulation in *Tmem16f* 292 cKO adapting neurons (unpaired mean difference brush: -11.4 [95% CI -19.2, -4.5],

vF: -6.4 [95% CI -8.5, -4.5]) as well as a reduction in spontaneous activity (Figure 4c-

- f, Figure 4—figure supplement 1a).
- In contrast, brush receptive field sizes and brush-evoked responses did not differ between control and *Tmem16f* cKO mice in non-adapting (inhibitory) neurons (Figure 4g-j), though an increase in vF-evoked responses in *Tmem16f* cKO animals was observed (mean difference 2.74 [95.00% Cl -0.074, 6.25], F(1; 110) = 7.014, P =0.0093) (Figure 4k, I). Spontaneous activity was also unaltered (Figure 4—figure supplement figure 1b).



301 302

#### Figure 4. Neonatal *Tmem16f* deletion in microglia decreases evoked activity, but increases

#### 303 brush receptive field size

- **a.** Brush receptive field sizes was increased for adapting neurons in *Tmem16f* cKO, mean difference
- 305 13.7 [95.00% CI 3.3, 27.2].
- 306 **b.** Overlay of receptive fields in **a**.
- 307 c. Brush evoked response was decreased for adapting neurons in Tmem16f cKO, mean difference -
- 308 11.4 [95% CI -19.2, -4.5].
- 309 d. Example firing trace of adapting cell from control and cKO animals to brush stimulation with raster310 plots underneath.
- 311 e. vF evoked activity was decreased for adapting neurons in *Tmem16* cKO, mean difference -6.4
- 312 [95% CI -8.5, -4.5].

f. Example firing trace of an adapting cell from control and cKO animals to vF stimulation with rasterplots underneath.

**g.** Brush receptive field sizes was unchanged for non-adapting neurons in *Tmem16f* cKO, mean

316 difference 5.53 [95.00% CI -15.91, 40.43].

317 h. Overlay of receptive fields in g.

318 i. Brush evoked response was unchanged for non-adapting neurons in *Tmem16f* cKO, mean

difference 8.62 [95.00% CI -5.78, 22.72].

320 j. Example firing trace of non-adapting cell from control and cKO animals to brush stimulation with321 raster plots underneath.

k. vF evoked activity was decreased for non-adapting neurons in *Tmem16f* cKO, mean difference
2.74 [95.00% CI -0.074, 6.25].

324 I. Example firing trace of a non-adapting cell from control and cKO animals to vF stimulation with325 raster plots underneath.

N-numbers as indicated. Black and colourful data points indicate females and males respectively.327

#### 328 Discussion

329 In summary, we report that dorsal horn microglia have a distinct phenotype during 330 the first postnatal week characterized by high phagocytic activity, which coincides 331 with postnatal engulfment of A-fibre terminals in the dorsal horn, supporting a role of 332 microglia in developmental dorsal horn remodelling. We further show that disruption 333 of microglial function by targeted deletion of *Tmem16f* during early postnatal life 334 impairs microglia mediated A-fibre refinement in the dorsal horn leading to long-term 335 changes in dorsal horn function and behaviour that persist into adulthood. Together, 336 our data suggests that microglia mediated refinement of A-fibres during the early 337 postnatal period is critical to both normal dorsal horn development and appropriate 338 spatial encoding of dynamic touch.

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In our study we looked at a subset of VGLUT1 positive A-fibres, which are predominantly low-threshold, myelinated A $\beta$  mechanoreceptors <sup>24,35</sup>, and primarily transduce innocuous mechanical stimulation of the skin, such as stroking and brushing. In adults, the superficial laminae is occupied by both A- and C-fibre projections, but the during the neonatal period A-fibre inputs dominate the superficial 345 dorsal horn, as nociceptive C-fibre inputs are weak and only start to strengthen between P5-P10 <sup>36,37</sup>. Functionally, this results in lower cutaneous mechanical 346 347 thresholds for activity in the superficial laminae, which together with immature local and descending inhibition leads to exaggerated reflex behaviour in neonates 5-7,38. 348 349 This early sensitivity to low threshold stimuli might be particularly important during 350 the neonatal period for maternal bonding, as skin to skin contact after birth was 351 shown to reduce neonatal stress and pain response and improve mother-infant interaction <sup>39</sup>. Therefore, appropriate A-fibre maturation might play an important role 352 353 in overall infant development.

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355 Microglia cells show the strongest increase in density over the first postnatal week 356 which coincides with a decrease in phagocytic activity. Hammond et al. 2019 357 revealed a specialised phagocytic population of microglia at P4/5 in the brain, while 358 our data suggests that the peak of microglial phagocytic activity might be later in the 359 spinal cord around P10. Moreover, dorsal horn microglial density observed in our 360 study appears to be higher than in various brain regions studied at comparable ages <sup>21,40</sup> (Table 1). Together, this suggests that dorsal horn microglia follow a different 361 362 developmental trajectory than brain microglia.

Paper	Brain area	Age	Density per µm <sup>3</sup>
Schwarz et al. 2012 <sup>21</sup>	Parietal cortex	P4	2.23 x 10 <sup>-7</sup>
	Amygdala	P4	4.77 x 10 <sup>-6</sup>
	CA1 of hippocampus	P4	4.88 x 10 <sup>-7</sup>
	Paraventricular nucleus	P4	1.37 x 10 <sup>-6</sup>
Perez-Pouchoulen et al. 2015 <sup>40</sup>	Cerebellum	P5	3 x 10 <sup>-6</sup>
Present study (Figure 1— figure supplement 1d)	Superficial dorsal horn	P3	5 x 10 <sup>-6</sup>

Table 1. Microglial density across different CNS regions (in rats). Estimated numbers
 from the graphs in respective paper, with male and female microglia counts pooled.

While microglial CD68 volume increased postnatally, the amount of microglial phagocytic cup numbers decreased. Phagocytic cups are relatively large structures involved in phagocytosing cell bodies <sup>41</sup>, and their reduction is consistent with the decline in apoptotic cell numbers postnatally. This shows that the postnatal increase in CD68 cannot be due to removal of apoptotic cell bodies but rather due to the refinement of neuronal connections.

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372 Dorsal horn microglia phagocytose A-fibre projections both in the superficial (laminae 373 I-II) and deeper (laminae III-IV) laminae during normal postnatal development. The 374 data here shows that most of the A-fibre engulfment occurs before P10 and 375 decreases thereafter. The overall amount of A-fibre engulfment is likely 376 underestimated, as reporter mice crossed with the Vglut1-Cre line used here only express tdT in a subset of A-fibres<sup>24</sup>. Further, the developmental upregulation of 377 Vglut1 expression <sup>26,27</sup> means that engulfment will be especially underestimated in 378 379 younger animals, as their A-fibres might be present but not yet expressing tdT. In the 380 brain, it has been shown that VGLUT1 protein expression at P0 is only 6.6% of the adult level, reaching 47% by P10 and 92% by P20<sup>27</sup>. Therefore, the fall in A-fibre 381 382 engulfment over the first postnatal week is likely much sharper in reality than Figure 383 1 g, h suggests.

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Although microglial CD68 volume peaks at P10, the engulfed VGLUT1-tdT volume seems to peak earlier at P7. While the percentage of lysosome volume occupied by VGLUT1-tdT is about 15% in the superficial laminae and 40% in the deep laminae at P7, by P10 it has become less than a quarter of that. It is possible that a slowing rate in engulfment of VGLUT1-tdT labelled fibres between P10 and P7 allows VGLUT1-

tdT to be degraded in the lysosomes at a faster rate than new VGLUT1-tdT material
is engulfed, thus causing a reduction of VGLUT1-tdT at P7 before the reduction in
lysosome volume follows suit by P10. It is also possible that developmental events at
P10 require additional microglial phagocytosis of materials other than A-fibres.

394

395 Neonatal *Tmem16f* cKO increased the number of Thy1-GFP positive VGLUT1 396 synapses in the superficial dorsal horn in adults, suggesting that *Tmem16f* function 397 is necessary for microglia mediated refinement of A-fibres.

However, increase in presynaptic terminals seems to predominantly affect lamina I, but not lamina III, even though developmental engulfment of A-fibres was observed both in LI-II and LIII-IV, suggesting that either only the engulfment of superficial synapses is dependent on *Tmem16f* or alternatively that superfluous synapses in the deeper laminae were more efficiently compensated and removed at later stages before animals reached adulthood.

The density of VGAT presynaptic terminals was also unaltered in *Tmem16f* cKO animals. This suggests that *Tmem16f* mediated engulfment is synapse specific. A recent study in the hippocampus suggested that complement mediated synapse elimination by microglia was targeted to VGLUT2, but not VGLUT1 or VGAT synapses instead <sup>42</sup>. Together with our data this suggest that developmental synapse elimination is both specific to synapse identity and location.

410 Consistent with the increase in dorsal horn VGLUT1, *Tmem16f* cKO increased brush 411 receptive field size in adapting/excitatory neurons, which mirrors the large receptive 412 field sizes in immature neonatal animals <sup>5</sup>. During the neonatal period A-fibre inputs 413 dominate the superficial dorsal horn <sup>36,37</sup> and hindpaw receptive fields are initially 414 large, reducing in size over the first two postnatal weeks <sup>5</sup>. Functionally, this results

in lower cutaneous mechanical thresholds, which together with immature local and
descending inhibition leads to exaggerated reflex behaviour in neonates <sup>5–7,38</sup>. Both
enlarged receptive field sizes and hypersensitivity were observed in Tmem16f cKO
animals. This is unlikely due to local disinhibition <sup>43,44</sup>, as spontaneous and evoked
activity of adapting WDR neurons were not increased, suggesting that the larger
receptive field sizes reflect a lack of primary afferent refinement instead.

421 Consistent with the increased A-fibre input and brush receptive field size, which likely 422 affects spatial summation of dynamic inputs, *Tmem16f* cKO increased behavioural 423 sensitivity to brush. However, counterintuitively, the increase in A-fibre projections 424 and behavioural sensitivity was accompanied by a reduction in evoked activity for 425 brush and vF as well as spontaneous activity in adapting neurons.

426 How does decreased evoked activity result in higher reflex sensitivity? One 427 possibility is that the activation pattern of WDR neurons may be more important than 428 firing rate per se for the coding of sensory information. Although the activity of 429 individual neurons is decreased, potentially more neurons are activated in total due 430 to the lack of A-fiber refinement. Therefore, the activation of WDR neurons on a 431 population level might be a better predictor of the spinal reflex response. A potential 432 mechanism underlying this effect is homeostatic synaptic downscaling, which 433 maintains homeostasis by reducing neuronal firing rate when the overall network activity is increased <sup>45</sup>. Our data shows that activity recorded from single neurons do 434 435 not always positively correlate with behavioural changes and need to be interpreted 436 with care.

437

438 In contrast to the adapting neurons, little change was observed for non-439 adapting/inhibitory neurons, apart from an increase in vF-evoked activity. This is

440 likely due to the low sample size, as fewer non-adapting/inhibitory neurons were 441 recorded. The increased firing rate of non-adapting/inhibitory neurons during vF-442 evoked activity would likely decrease the activity of adapting/excitatory neurons and 443 thus contribute to the decreased overall activity observed in cKO animals.

444 We propose therefore that disrupting microglial function with *Tmem16f* cKO causes 445 an increase in VGLUT1 positive A-fibre terminals in adult animals, potentially due to 446 failed pruning of A-fibres in the neonatal period. This excess of peripheral input might 447 preferentially target excitatory neurons which leads to larger receptive field sizes 448 thereof, and potentially activates larger numbers of excitatory neurons upon 449 peripheral stimulation. In compensation, through homeostatic scaling or increased A-450 fibre input onto non-adapting/inhibitory neurons, inhibitory activity onto excitatory 451 neurons is increased, decreasing the firing rate of individual neurons, but not 452 necessarily the total output as a population. This is also consistent with increased 453 behavioural sensitivity in *Tmem16f* cKO animals.

454

455 Limitations

456 Several caveats need to be considered when interpreting the data. In our 457 experiments we used a CNS-wide KO of microglial Tmem16f. Therefore, we cannot 458 rule out cortical contributions to the observed changes in the spinal cord - for 459 example, top-down facilitation of inhibitory inputs or inhibition of excitatory inputs 460 could also explain the decreased firing rate in adapting and non-adapting neurons in 461 Tmem16f cKO animals. However, a purely top-down mediated overall inhibition of 462 dorsal horn activity would not explain the increase in receptive field size following 463 *Tmem6f* 1 cKO, though it could additionally contribute to the changes observed.

464 Therefore, at least part of the observed changes must be mediated by changes in465 peripheral afferent input to the dorsal horn.

Furthermore, it is possible that *Tmem16f* cKO will also affect the engulfment and removal of apoptotic cells by microglia, which contribute to the behavioural and functional changes seen in adulthood. However, the mechanisms involved in engulfing whole cell bodies vs synaptic material are likely distinct (in our study CD68 volume was not correlated with apoptotic clearance), and therefore *Tmem16f* cKO might affect microglial pruning without affecting the phagocytosis of apoptotic cells.

472 We used VGLUT1 as a marker for presynaptic boutons of A-fibres as well as the 473 driver of tdT expression. However, it has been shown that cortico-spinal projections also contribute to VGLUT1 synapses throughout the dorsal horn <sup>33</sup>. Therefore, we 474 475 cannot exclude that the behavioural and functional effects seen in the Tmem16f cKO 476 animals are due to disruption of cortico-spinal refinement. However, during early 477 postnatal development, corticospinal projections do not enter in the grey matter 478 lumbar segment until P7, while peripheral A-fibre afferents reach the lumbar dorsal horn by embryonic day (E)14<sup>46,47</sup>. Therefore, we can conclude that at least prior to 479 480 P7, where the majority of VGLUT1-tdT engulfment was observed, the engulfment of 481 VGLUT1-tdT labelled fibres are indeed afferent A-fibres.

An important open question is what determines the engulfment or survival of synapses and fibres in the dorsal horn. It is known that microglia can sense and respond to neuronal activity <sup>48–51</sup>. For example, microglia increase process motility both to neuronal hyper- and hypoactivity <sup>49</sup>, and in the brain, microglial engulfment seems to preferentially target weak synapses <sup>12,13</sup>. Given that perturbation of neuronal activity can indeed impair A-fibre refinement <sup>8,9</sup>, neuronal activity seems a likely signal for microglial engulfment in the dorsal horn. Several molecules

expressed by neurons and microglia have been identified as "eat me" and "keep me"
signals <sup>12,13,52,53</sup>, among which members of the complement cascade, e.g. C1q, have
been shown to tag synapses for microglial engulfment both in the brain and the
spinal cord <sup>12,17</sup>. Whether C1q also plays a role in dorsal horn remodeling remains to
be investigated.

494 Conclusions

In summary, we have shown that dorsal horn microglia phagocytose A-fibres during normal postnatal development and that disruption of microglial function can lead to long-term structural and functional changes in the dorsal horn and behavioural changes towards dynamic touch. This has important implications for perinatal care of infants.

500 Low-threshold stimuli such as skin-to-skin contact have been shown to reduce infant stress and improve infant-mother interaction and maternal bonding <sup>39</sup>. Therefore, 501 502 appropriate maturation of A-fibres might play an important role in overall infant 503 development. In addition, previous research have shown that peripheral injury during 504 the first postnatal week (but not after P10) cause hyperalgesia upon re-injury in 505 adulthood, and that this could be prevented with intrathecal minocycline injection, which is a non-specific microglial inhibitor <sup>54,55</sup>. It is tempting to speculate that injury 506 507 alters microglial pruning of A-fibres as described here which could underlie part of 508 the long-term changes seen following injury. Thus, the role of microglia in normal 509 and aberrant neonatal development could be investigated for therapeutic potential in 510 perinatal care.

511

### 512 Materials and Methods

Key Resources Table				
Reagent type (species) or resource	Designation	Source or reference	Identifiers	Additional informatio n
Genetic reagent ( <i>M.</i> <i>musculus</i> )	Ai9; <i>R26<sup>LSL-Ai9/+</sup></i>	Jackson Laboratory	Stock # 007909 RRID:IMSR_JAX:00 7909	PMID: 20023653
Genetic reagent ( <i>M.</i> <i>musculus</i> )	Ai9; <i>R26<sup>LSL-Ai9/+</sup></i>	Jackson Laboratory stock	Stock # 007905 RRID:IMSR_JAX:00 7905	PMID: 20023653
Genetic reagent ( <i>M.</i> <i>musculus</i> )	<i>Slc17a7</i> -IRES2- Cre; <i>Vglut1<sup>Cre/+</sup></i>	Jackson Laboratory stock	Stock # 023527 RRID:IMSR_JAX:02 3527	PMID: 25071457
Genetic reagent ( <i>M.</i> <i>musculus</i> )	Cx3cr1- CreERT2-YFP; Cx3cr1 <sup>CreER/+</sup>	Jackson Laboratory stock	Stock # 021160 RRID:IMSR_JAX:02 1160	PMID: 35045285
Genetic reagent ( <i>M.</i> <i>musculus</i> )	Thy 1-EGFP-M; Thy 1 <sup>eGFP</sup>	Jackson Laboratory stock	Stock # 007788 RRID:IMSR_JAX:00 7788	PMID: 11086982
Genetic reagent ( <i>M. musculus</i> )	<i>Tmem16f-</i> floxed; <i>Tmem16f<sup>fl/fl</sup></i>	PMID: 27332874		Dr Paul Heppenst all (SISSA, Trieste, Italy)
Antibody	anti-Caspase-3 (Rabbit monoclonal)	Cell Signalling	Cat# Asp175, 5A1E	IF (1:100)
Antibody	anti-CD68 (Mouse monoclonal)	BIO-RAD	Cat# MCA341R	IF(1:500)
Antibody	anti-CD68 (Rabbit polyclonal)	Abcam	Cat# ab125212	IF(1:500)

Antibody	anti-GFAP (Mouse, monoclonal)	Sigma	Cat# G3893	IF(1:2000)
Antibody	anti-Iba1 (Rabit, polyclonal)	Wako	Cat# 019-19741	IF(1:1000)
Antibody	anti-Iba1 (Goat, polyclonal)	Abcam	Cat# ab5076	IF(1:1000)
Antibody	anti-NeuN (Mouse, monoclonal)	Millipore	Cat# MAB377	IF(1:2000)
Antibody	anti-VGAT (Mouse, monoclonal)	Synaptic Systems	Cat# 131011	IF(1:2000)
Antibody	anti-VGAT (Rabbit, polyclonal)	Synaptic Systems	Cat# 131002	IF(1:2000)
Antibody	anti-VGLUT2 (Guinea pig, polyclonal)	Millipore	Cat# AB2251-I	IF(1:2000)
Antibody	anti- Synaptophysin (Mouse, monoclonal)	Abcam	Cat# Ab8049	IF(1:2000)
Antibody	anti- Synaptophysin (Guinea pig, polyclonal)	Synaptic Systems	Cat# 101004	IF(1:2000)
Antibody	anti-goat Cy2 (Donkey polyclonal)	Jackson ImmunoResea rch	Cat# 705-225-147	IF(1:500)
Antibody	anti-goat Cy3 (Donkey polyclonal)	Jackson ImmunoResea rch	Cat# 705-165-147	IF(1:500)
Antibody	anti-goat Cy5 (Donkey polyclonal)	Jackson ImmunoResea rch	Cat# 705-175-147 1	IF(1:500)

Antibody	anti-goat Alexa 488 (Donkey polyclonal)	Jackson ImmunoResea rch	Cat# 705-545-003	IF(1:500)
Antibody	anti-guinea pig Cy3 (Donkey polyclonal)	Jackson ImmunoResea rch	Cat# 706-165-148	IF(1:500)
Antibody	anti-guinea pig Alexa 647 (Donkey polyclonal)	Millipore	Cat# AP193SA6	IF(1:500)
Antibody	anti-mouse Cy2 Jackson (Donkey polyclonal)	Jackson ImmunoResea rch	Cat# 715-225-150	IF(1:500)
Antibody	anti-mouse Cy3 Jackson (Donkey polyclonal)	Jackson ImmunoResea rch	Cat# 715-165-150	IF(1:500)
Antibody	anti-mouse Cy5 (Donkey polyclonal)	Jackson ImmunoResea rch	Cat# 715-175-150	IF(1:500)
Antibody	anti-rabbit Cy2 (Donkey polyclonal)	Jackson ImmunoResea rch	Cat# 711-225-152	IF(1:500)
Antibody	anti-rabbit Alexa 594 (Donkey polyclonal)	Jackson ImmunoResea rch	Cat# 711-585-152	IF(1:500)
Antibody	anti-rabbit Alexa 647 (Donkey polyclonal)	Millipore	Cat# AP182SA6	IF(1:500)
Antibody	anti-rabbit- biotinylated (Horse polyclonal)	Vector	Cat# BA11-00	IF(1:500)
peptide, recombin ant protein	Streptavidin-594	Life Technologies	Cat# S32356	IF(1:500)

Software, algorithm	ImageJ; FIJI	PMID:229308 34, PMID:291871 65	RRID:SCR_003070	
Software, algorithm	Volocity	PerkinElmer	RRID:SCR_002668	
Software, algorithm	LabChart 7	ADInstruments		
Software, algorithm	Graphpad prism	GraphPad Software, Inc.	RRID:SCR_002798	
Software, algorithm	Estmationstats. com (web application)	PMID: 31217592	RRID:SCR_018321	Free web applicatio n
other	Carbostar-1 (micro- electrode)	Kation Scientific		

513

#### 514 Animals

- 515 Sprague Dawley rats of both sexes were used for experiments in Figure 1—figure
- 516 supplement 1d-h and Figure 1—figure supplement 2. Transgenic mice on C57BL/6J
- 517 background of both sexes were used in all other experiments.
- 518 Experiments used the following transgenic mouse lines:
- 1. Ai9 / Rosa26-CAG::loxP-STOP-loxP-tdTomato-WPRE (Jackson Laboratory
- 520 stock 007909 & 007905)
- 521 2. *Slc17a7*-IRES2-Cre (Jackson Laboratory stock 023527)
- 522 3. *Cx3cr1*-CreERT2-YFP (Jackson Laboratory stock 021160)
- 523 4. *Thy1*-EGFP-M (Jackson Laboratory stock 007788)
- 524 5. *Tmem16f*-floxed (flx) animals (Batti et al., 2016)
- 525 For visualisation of A-fibres, Slc17a7-IRES2-Cre (Vglut1-Cre) males (JAX stock no.
- 526 023527) were crossed with Ai9 females (JAX stock no. 007909) to obtain animals

527 that expressed the tdTomato fluorophore under the *Vglut1* promoter (*Vglut1*<sup>Cre/+</sup>; 528  $R26^{LSL-Ai9/+}$ ).

529 To generate tamoxifen inducible microglia-specific *Tmem16f* knock-out mice 530 (*Tmem16f* cKO), Cx3cr1-CreER-YFP (JAX stock no. 021160) mice were crossed to 531 *Tmem16f*-flx animals (generated by P. Heppenstall, see Batti et al. 2016), as well as 532 Ai9 (JAX stock no. 007905) and *Thy1*-EGFP-M (JAX stock no. 007788).

- Experimental animals were heterozygous for *Cx3cr1*-CreER-YFP ( $^{CreER/+}$ ), homozygous for mutant conditional allele *Tmem16f*-flx ( $^{fl/fl}$ ), and carrying Ai9 ( $R26^{LSL-}$ <sup>Ai9</sup>) and *Thy1*-eGFP ( $^{eGFP}$ ) alleles (zygocity was not determined for  $R26^{LSL-Ai9}$  and *Thy1*<sup>eGFP</sup>). This produced the following genotype:  $Cx3cr1^{CreER/+}$ ; *Tmem16f*<sup>fl/fl</sup>;  $R26^{LSL-}$ <sup>Ai9</sup>; *Thy1*<sup>eGFP</sup>. Control animals were homozygous for the wild type *Tmem16f* allele: *Cx3cr1*<sup>CreER/+</sup>; *Tmem16f*<sup>+/+</sup>;  $R26^{LSL-Ai9}$ ; *Thy1*<sup>eGFP</sup>.
- 539 Both females and males were used. No sex differences were expected and animals 540 of both sexes were pooled together for analysis, but data points are presented as 541 black (female) or red/magenta/blue (male) to indicate the sexes. Numbers of animals 542 used for each experiment are indicated in the figures. For a table with detailed 543 species, ages, sexes, and numbers of animals used in each experiment, please see 544 supplementary Table 1. All procedures were carried out in accordance with the 545 guidelines of the UK Animals (Scientific Procedures) Act 1986 and subsequent 546 amendments.

547

#### 548 **Drugs**

549 4-hydroxytamoxifen (4-HT) was dissolved at 1mg/ml in corn oil, and 50µl was 550 injected intragastrically per pup daily on three consecutive days from P1-3, following

a previously described protocol  ${}^{56}$ . Both control and experimental animals (*Tmem16*<sup>+/+</sup> and *Tmem16f*<sup>#/#</sup> respectively, see animals section above) received 4-HT injections to control for any effects of 4-HT itself. The dam was given a protein enriched diet a few days before and following delivery to aid milk production and pup survival.

#### 556 *Immunohistochemistry*

557 Animals were overdosed with pentobarbital and transcardially perfused with saline 558 followed by ice-cold 10% formalin. The sciatic nerve was exposed and traced to 559 locate L4 & L5 dorsal root ganglia (DRG) and the corresponding region of the lumbar 560 spinal cord was dissected and post-fixed in 10% formalin overnight, followed by 561 immersion in 30% sucrose until they sank. 50µm free-floating spinal cord sections 562 were cut on the microtome with every 2nd section collected.

563 Tissue sections were washed 3 x 10 min in PBS and then incubated in blocking 564 solution (10% donkey serum, 0.2% Triton X-100 in PBS) for 2.5h at room 565 temperature. The sections were then incubated with primary antibodies at 4°C 566 overnight followed by secondary antibodies at room temperature for 2h, both diluted 567 in 3% blocking solution (3% donkey serum, 0.2% Triton X-100 in PBS) (for list of antibodies and their respective concentrations used, see Key Resources Table). 568 569 Samples were mounted in Fluoromount Aqueous Mounting Medium (Sigma) or ProLong<sup>™</sup> Diamond Antifade Mountant (Thermo Fischer), if the tissue contained 570 571 endogenous fluorophores.

572

#### 573 Image acquisition and analysis

574 Confocal z-stacks were taken with a Zeiss LSM880 confocal microscope or 575 Yokogawa CSU22 spinning disk microscope using a 20x water immersion objective 576 (NA 1.0) for imaging of A-fibres and 63x oil immersion objective (NA 1.4) followed by 577 analysis in Fiji software. Details on microscope settings can be found in 578 supplementary Table 1 and in the metadata of example images online at 579 https://github.com/Yajing826/A-fibre-engulfment. Only intact sections with an even 580 stain were analysed, and at least 6 sections were imaged and analysed per animal 581 to reduce variability for all figures (apart from S3, where 1-3 sections were analysed 582 per animal).

583

584 Cell counts and phagocytic cup counts were performed manually on confocal images 585 in Fiji or Volocity, with the experimenter blinded to the age groups. Phagocytic cups 586 were defined as any rounded structure at the end of a microglial process.

587 A-fibre engulfment by microglia and synapse density were analysed with automated 588 batch processing in Fiji using the 3D-ROI manager plugin and custom written macros <sup>57–60</sup>. For A-fibre engulfment, each of the channels containing staining for microglia, 589 590 lysosomes, or A-fibres were binarized and the volume of their overlap measured. For 591 synapse density measures, the channel containing synaptic stain was binarized and 592 segmented, following which volume and object numbers were recorded. Macro 593 scripts the analysis available online for automated are at 594 https://github.com/Yajing826/A-fibre-engulfment.

595

#### 596 **Behaviour**

597 Behavioural testing was carried out on adult mice of both sexes between 3-4 months 598 old, with the experimenter blinded to animal genotype/treatment. Animals were 599 placed on a mesh platform (Ugo Basile) within individual transparent plastic 600 chambers (6cm x 6cm x 12cm) for sensory testing of the plantar surface of the hind 601 paw. Habituation and testing happened over five consecutive days. Animals were 602 habituated to the testing environment for 1h per day on the first two days within 603 individual plexicon chambers on the mesh platform. On the remaining days, animals 604 were habituated for 30 min before being tested on each day. Brush response and 605 von-Frey (vF) threshold were determined on the 3rd day, while repeated vF 606 response testing was spread over the remaining 2 days to avoid sensitisation. 607 Number of withdrawal reflexes were scored in each case, where only a rapid paw 608 lifting was scored as a reflex. Animals were allowed to rest at least 20 seconds 609 between each stimulus. For brush response, a fine brush (Pro Arte, series 007, size 610 2) was moved over the plantar surface of the hind paw from heel to toe over a 2 second period <sup>61</sup>. This was repeated five times and the number of withdrawal 611 612 reflexes out of five was recorded. vF threshold was assessed using the simplified up and down method <sup>62</sup>. Filaments were aimed at the region between the foot pads. 613 614 Force was applied until the filament bent, and held in place for 2 seconds.

To generate a response curve to vF stimulation, repeated vF response was recorded by applying each of filaments no. 3 - 7 (0.04g - 0.6g) five times on the plantar surface, directed at the region between the foot pads. The sequence of vF filaments was randomised. Number of withdrawal reflexes out of five times was recorded.

619

#### 620 In vivo extracellular recording

Animals subjected to behavioural testing were reused in electrophysiological recordings. Experimenter was blinded to animal genotype/treatment. All recordings were performed on adult mice (3-4 months old) of both sexes in the deep dorsal horn. Cells were not recorded beyond 550 µm depth from the surface of the spinal cord (see Figure 4—figure supplement 1c for depth of all recorded neurons). 3-5 mice were used per sex and treatment group. All experiments were carried out by the same experimenter to ensure consistency.

#### 628 Animal preparation

Mice were anaesthetised with intraperitoneal urethane injection (10% in saline, 1.5g/kg). 100µl of 0.6 g/ml atropine and 200µl saline were injected subcutaneously to respectively counteract the mucus-driving side effect of urethane and to prevent dehydration. The animal was constantly monitored for depth of anaesthesia throughout the experiment and supplemented with 50µl (5µg) urethane as needed. 200µl of saline was supplemented every 2 hours. Body temperature of the animal was kept close to 37°C with a heating pad throughout.

After cessation of reflexes, a tracheotomy was performed and a short plastic tube of about 1cm inserted to aid free breathing of the animal. The animal was then transferred onto a stereotactic frame and fixed with ear and hip bars. A laminectomy was carried out at vertebral level T13-L1 which corresponds to the spinal segments L4-L5 underneath. The spinal column was clamped for stability, the dura was removed, and the exposed spinal cord was covered with mineral oil to prevent drying.

#### 643 <u>Single unit extracellular recordings</u>

644 A carbon micro-electrode (Carbostar-1, Kation Scientific) was lowered with a 645 motorised manipulator (Scientifica) into the exposed spinal cord at a constant speed. 646 A reference electrode was inserted into the back muscle close to the laminectomy for 647 differential recording. Recorded neural activity was amplified 2000 times, and filtered 648 for signals between 1kHz-10kHz (NL104 amplifier and NL125/6 band-pass filter 649 modules from NeuroLog Digitimer). The signal was sampled at 20kHz and digitised 650 using Powerlab 4/30 (ADInstruments). The trace was recorded and analysed in the 651 software LabChart 7 (ADInstruments).

To isolate single neurons, the plantar surface of the animal's hind paw was gently continuously stroked as a searching signal, while the electrode is being lowered through the dorsal horn of the spinal cord. Once a cell has been identified by equal amplitude of the spikes recorded, the brush receptive field of the cell was mapped out by a fine brush (Pro Arte, series 202, size 1, brush tip cut short to 7mm length x 1mm width).

Spontaneous activity was recorded for 10 min before and 5 min after stimulation. To record brush and von-Frey (vF) fibre evoked activity from single neurons, each stimulus was manually applied for 2 seconds over the receptive field of the cell and repeated 3 times, with a minimum of 10 second interval in between. (vF filament strength were as follows: 1 = 0.008g, 2 = 0.02g, 3 = 0.04, 4 = 0.07g, 5 = 0.16g, 6 =0.40g, 7 = 0.60g, 8 = 1g, 9 = 1.7, 10 = 2g).

664 Cells were not recorded if they exhibited very high spontaneous firing rates that did 665 not allow evoked activity to be clearly distinguished from spontaneous activity. Only 666 wide dynamic range neurons responding both to brush and pinch stimulation were 667 recorded. Animals were euthanised at the end of the experiment and the spinal cord

was collected in neutral buffered 10% formalin (overnight) followed by 30% sucrosesolution for re-use in immunohistochemistry.

#### 670 Analysis & cell type categorisation

Analysis was carried out in the LabChart 7 software (ADInstruments). For spontaneous activity, firing rate was analysed over a 10min window prior to applying any stimuli. Firing rate for evoked responses were analysed over the first second of the stimulus duration and averaged over three trials.

675 Cells were divided into adapting and non-adapting groups based on their firing 676 properties towards a threshold vF-stimulus  $^{32}$ . This threshold vF was defined as the 677 first vF filament that evokes a firing rate of 10Hz or more. The response within the 678 first second of vF application was

analysed to calculate an adaptive ratio R, which was defined as

$$R = \frac{Number \ of \ spikes \ fired \ between \ 0.5 - 1 \ sec}{Number \ of \ spikes \ fired \ between \ 0 - 0.5 \ sec}$$

If a cell adapts rapidly to stimulation, one would expect R to be close to zero, as barely any spikes should be fired between 0.5 - 1 sec, however, if a cell is nonadapting and firing continuously, one would expect R to be close to 1. To decide the boundary between adapting and non-adapting cells we used k-means cluster analysis, which sorted the values into two groups that is equivalent to a boundary at R = 0.33. For the k-means clustering we included cells from a previous experiment.

#### 686 Statistical Analysis

Estimation statistics for the 95% confidence intervals (95% CI) of the mean difference were calculated on estimationstats.com <sup>63</sup> using 5000 samples of biascorrected and accelerated bootstrapping. As bootstrapping is less accurate for small

samples sizes <sup>64</sup>, confidence intervals were only calculated for samples with N≥5. For Figure 1—figure supplement 1d, h and Figure 1—figure supplement 2 where N=4, one-way ANOVA was used with post-hoc comparisons carried out using Dunnett's method.

Additionally, conventional null-hypothesis significance testing was carried out on estimationstats.com and GraphPad Prism 6 for all comparisons (significance level was set at  $\alpha$ =0.05), which are listed in the Appendix 1—table 1.

Data are presented as mean ± SD in all figures. Where applicable, the effect size is presented as 95% CI of the mean difference on a separate but aligned axis. The mean difference is plotted as a dot on the background of its probability distribution, and the 95% confidence interval is indicated by the ends of the error bar. All values in text and figures are given with two decimals or rounded to two significant figures. N-numbers are as indicated in figures. For a comprehensive list with exact statistical values and analyses, see Appendix 1—table 1.

704

#### 705 Data availability

706 All data generated or analysed during this study are included in the manuscript and 707 supporting files. Source files for all data available are at: 708 https://www.ebi.ac.uk/biostudies/ under the accession number S-BSST609. Macro 709 scripts for automated analysis in Fiji are available online at 710 https://github.com/Yajing826/A-fibre-engulfment.

711

712

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#### 716 Author Contributions

YX and SB performed the experiments, YX, SK, MF and SB designed the study, analysed and interpreted data and co-wrote the manuscript. AC, QH and RRJ developed the mouse model and prepared tissue for analysis. MS and PH developed the mouse model.

#### 721 Competing Interest Statement

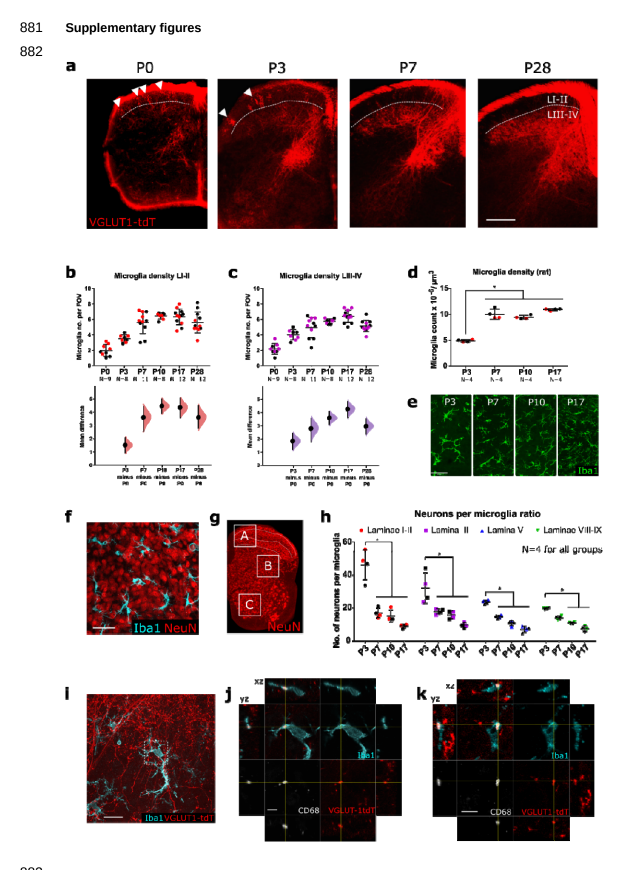
The authors declare no competing interests.

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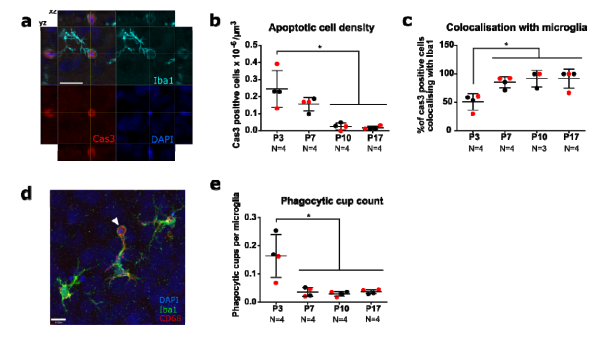
a. Representative images showing an increase in tdT-expression in the dorsal horn with age. Dashed
 white line indicates border between lamina II and lamina III, white arrow heads indicate tdT-fibres in
 the superficial laminae I-II. Scale bar = 200µm.

b. Microglial density increases over the postnatal period in both LI-II. P3 vs. P0: mean difference 1.53
[95% CI 0.97, 2.08]; P7 vs. P0: mean difference 3.61 [95% CI 2.58, 4.42]; P10 vs. P0: mean difference 4.46 [95% CI 3.91, 4.98]; P17 vs. P0: mean difference 4.35 [95% CI 3.60, 5.06]; P28 vs.
P0: mean difference 3.61 [95% CI 2.79, 4.49]. Field of view (FOV) = 245µm x 65µm.

**c.** Microglial density increases over the postnatal period in LIII-IV. P3 vs. P0: mean difference 1.83 [95% CI 1.21, 2.42]; P7 vs. P0: mean difference 2.79 [95% CI 1.80, 3.54]; P10 vs. P0: mean difference 3.58 [95% CI 3.09, 4.06]; P17 vs. P0: mean difference 4.24 [95% CI 3.60, 4.81]; P28 vs. P0: mean difference 2.96 [95% CI 2.35, 3.51]. Field of view (FOV) =  $245 \mu m \times 65 \mu m$ .

- 896 **d.** Microglia density in rats increases significantly between P3 and P7, with P < 0:0001 for all ages compared to P3.
- 898 e. Representative images of microglia cells in the dorsal horn (LI-III) stained for Iba1 (green) at
   899 postnatal days (P)3, P7, P10, P17. Scale bar = 50 μm.
- 900 f. Maximum projected confocal images from the dorsal horn (LI-III) at P3 with representative staining
   901 for microglia (Iba1, cyan) and neurons (NeuN, red). Scale bar = 50µm.
- 902 g. P7 spinal cord hemisection stained with neuronal marker NeuN (red), white boxes A, B, C indicate
   903 the approximate area of laminae I-III, lamina V, and laminae VIII-IX used for analysis.
- 904 h. Ratio of neurons per microglia significantly decreases with age in laminae I-II, lamina III, lamina V,
   905 and laminae VIII-IX.
- 906 i. Representative confocal image of microglial A-fibre engulfment in the spinal dorsal horn of P3
   907 VGLUT1-tdt mice, stained for microglia (Iba1, cyan), microglial lysosomes (CD68, white) and
   908 endogenously fluorescent A-fibres (tdTomato, red). Scale bar = 20µm. White inset box show location
- 909 of higher magnification panels in **i.**
- 910 j. High magnification image of microglial A-fibre engulfment in h stained for microglia (Iba1, cyan),
- 911 microglial lysosomes (CD68, white) and endogenously fluorescent A-fibres (tdTomato, red). Cross-
- hairs show position of the xz and yz side-view panels. Scale bar =  $5\mu$ m.
- 913 **k.** Further example of microglial A-fibre engulfment, colours and scale as indicated in **i**.
- 914 N-numbers as indicated. Black and colourful data points indicate females and males respectively.





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#### 919 Figure 1—figure supplement 2. Apoptotic cell clearance over the postnatal period

a. Representative confocal image from the P3 dorsal horn of a microglia cell engulfing an apoptotic cell with a phagocytic cup. Cross-hairs show position of the xz and yz side-view panels. Microglial cell body is outside of the visible xy-plane. Image shows representative staining for Iba1 (cyan), Cas3
(red), DAPI (blue) at P3. Scale bar = 20µm.

b. Apoptotic cell numbers decrease significantly by P10, with P < 0.001 for P10 and P17 compared to</li>
P3.

926 c. The percentage of Cas3 positive cells colocalising with microglia significantly increases with age,
 927 with P < 0.05 for all ages compared to P3.</li>

d. Representative image of microglia cell in the dorsal horn with phagocytic cup indicated by white
 arrowhead. Image shows representative staining for Iba1 (green), CD68 (red), DAPI (blue) at P3.
 Scale bar = 10µm.

e. Phagocytic cup count per microglia decreases significantly between P3 and P7, (P < 0.01 for all ages compared to P3).</li>

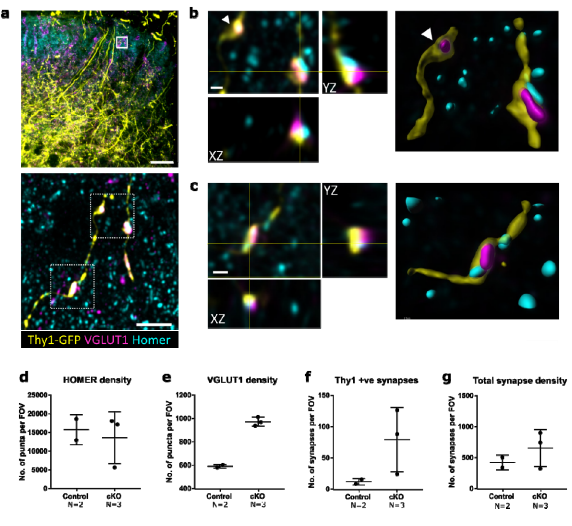
933 N-numbers as indicated. Black and red data points indicate females and males respectively.

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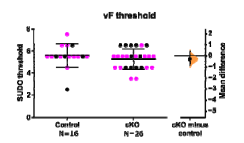
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# Figure 2—figure supplement 1. Superficial A-fibres form synapses with pre- and post-synaptic densities.

- **a.** TOP: Low magnification image, with white box indicating the location of bottom panel image. Scale
- 944 bar = 50µm. Bottom: Representative maximum projection of super-resolution image from the
- 945 superficial dorsal horn laminae showing colocalization of Thy1-GFP labelled A-fibre (yellow) with
- presynaptic VGLUT1 (magenta) and postsynaptic HOMER (cyan). Scale bar = 5µm.
- 947 **b, c.** Super-resolution images of boxed areas in **a** (bottom panel) with xy, xz, and yz views showing
- 948 colocalization of Thy1-GFP labelled A-fibre (yellow), presynaptic VGLUT1 (magenta), and
- 949 postsynaptic HOMER (cyan), as well as surface rendered view on the right. White arrowhead in b
- 950 indicate a Thy1-GFP and VGLUT1 positive presynaptic bouton without postsynaptic HOMER. Scale
   951 bars = 1µm.
- 952 **d.** Homer density does not seem affected in *Tmem16f* cKO animals.
- 953 e. VGLUT1 density seems higher in *Tmem16f* cKO animals, consistent with results in Fig. 3.
- 954 f. Thy1-positive synapse density (as defined by colocalization of Thy1-GFP, VGLUT1, and HOMER)
   955 seems to increase in *Tmem16f* cKO animals.
- 956 **g.** Total synapse number (as defined by colocalization of VGLUT1 and HOMER puncta) seems
- 957 unchanged in *Tmem16f* cKO animals.
- 958 FOV=  $94\mu m \times 94\mu m$  for **d-g**. N-numbers as indicated.



#### 960 Figure 3—figure supplement 1. Tmem16f cKO does not change vF reflex threshold

961 Unpaired mean difference -0.29 [95% CI -0.85, 0.39]. N-numbers as indicated. Black and magenta
 962 data points indicate females and males respectively.

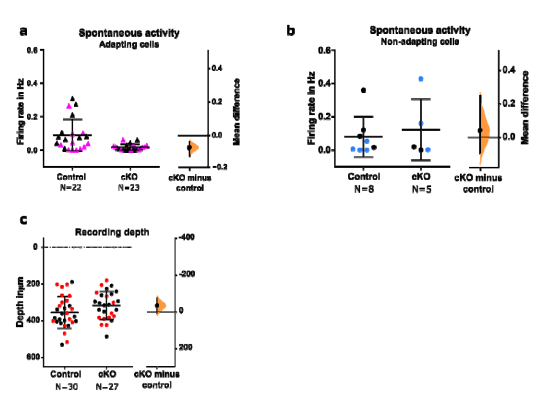
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## 968 Figure 4—figure supplement 1. Recording depth and spontaneous activity in control and

#### 969 *Tmem16f* cKO animals.

a Spontaneous activity decreased in adapting WDR neurons from *Tmem16f* cKO animals, unpaired
 mean difference -0.072 [95% CI -0.121, -0.039].

972 b. Spontaneous activity was unchanged in non-adapting WDR neurons from *Tmem16* cKO animals,
973 unpaired mean difference 0.042 [95.00% CI -0.092, 0.25].

974 c. Recording depth of all neurons in control and *Tmem16* cKO animals, unpaired mean difference -

- 975 37.24 [95.00% CI -78.02, 5.73].
- 976 N-numbers as indicated. Black and coloured data points indicate females and males.
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