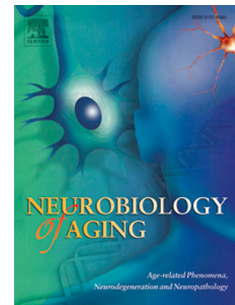


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Age-dependent changes in clock neuron structural plasticity and excitability are associated with a decrease in circadian output behaviour and sleep

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1   **Age-dependent changes in clock neuron structural plasticity and excitability are associated with a**  
2   **decrease in circadian output behaviour and sleep**

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8

9   **Abstract**

10   Ageing has significant effects on circadian behaviour across a wide variety of species, but the  
11   underlying mechanisms are poorly understood. Previous work has demonstrated the age-dependent  
12   decline in behavioural output in the model organism *Drosophila*. We demonstrate this age-  
13   dependent decline in circadian output is combined with changes in daily activity of *Drosophila*.

14   Ageing also has a large impact on sleep behaviour, significantly increasing sleep duration whilst  
15   reducing latency. We used electrophysiology to record from large ventral lateral neurons (l-LNv) of  
16   the *Drosophila* circadian clock, finding a significant decrease in input resistance with age, but no  
17   significant changes in spontaneous electrical activity or membrane potential. We propose this  
18   change contributes to observed behavioural and sleep changes in light-dark conditions. We also  
19   demonstrate a reduction in the daily plasticity of the architecture of the small ventral lateral neurons  
20   (s-LNv), likely underlying the reduction in circadian rhythmicity during ageing. These results provide  
21   further insights into the effect of ageing on circadian biology, demonstrating age-related changes in  
22   electrical activity in conjunction with the decline in behavioural outputs.

23

24   **1. Introduction**

25 Circadian rhythms describe the near 24-hour cycle in behaviour and physiology, driven by the  
26 circadian clock, that allow organisms to anticipate daily changes in their environment. Circadian  
27 clocks in animals are fundamental biological components responsible for the control of large aspects  
28 of physiology and behaviour, ranging from the sleep-wake cycle to rhythms in blood pressure  
29 (Roenneberg and Merrow, 2016). Remarkably, the fundamental molecular basis of the intracellular  
30 clock is well conserved from *Drosophila* to mice and humans (Allada and Chung, 2010). The health  
31 consequences of circadian misalignment as a result of our modern lifestyles are dramatic, with links  
32 to cancer, depression and sleep disorders (Menet and Rosbash, 2011; Samuelsson et al., 2018; West  
33 and Bechtold, 2015). With increasing human lifespans and an ageing population, understanding how  
34 circadian rhythms change during the ageing process is of growing interest and medical relevance,  
35 with the population aged over 60 years old set to more than double by 2050 (UN, 2015).

36 It is well established that elderly individuals have increasing difficulties sleeping at night and have  
37 and increase in daytime sleep episodes combined with generally going to sleep and waking up earlier  
38 (Kondratova and Kondratov, 2012). The daily cycles of hormone levels, body temperature and the  
39 sleep-wake cycle, are modified with age in humans causing disruption in behaviour, and resultant  
40 reduction in the strength of the clock (Hofman and Swaab, 2006). Furthermore, circadian sleep-wake  
41 disorders are more prevalent in older individuals (Kim and Duffy, 2018).

42 Using *Drosophila* offers numerous advantages for investigating how ageing affects the circadian  
43 clock, not least the strong history of circadian research in the model organism, genetic tractability,  
44 short lifespan (50-80 days), rapid generation time as well as clearly defined and manipulatable  
45 neural circuits. Genetic analysis of circadian behaviours has identified genes involved in generating  
46 rhythmic transcription translation feedback loops comprising the molecular clock of *Drosophila*  
47 (Allada and Chung, 2010; Hardin, 2011; Tataroglu and Emery, 2015), which in turn control a wide-  
48 range of physiological and cellular responses, likely through rhythmic control of output genes.

49 The *Drosophila* central clock consists of 150 dispersed but connected circadian neurons that are  
50 classified by their anatomical location, projection pattern and the expression of clock genes (Peschel  
51 and Helfrich-Förster, 2011). They function as a network to drive rhythmic behaviour (Top and Young,  
52 2017). Examples of outputs from the molecular clock are the circadian remodelling of the projections  
53 from the s-LNv clock neurons to the dorsal protocerebrum (Fernández et al., 2008) and circadian  
54 modulation of the firing frequency and membrane potential of clock neurons (Cao and Nitabach,  
55 2008; Flourakis et al., 2015; Sheeba et al., 2007). Under laboratory conditions using a 12:12 hr  
56 light:dark (LD) cycle, *Drosophila* display morning and evening peaks in locomotor activity with  
57 anticipation activity prior to the transitions of light-on and lights-off, with constant darkness (DD)  
58 resulting in free-running activity with a period of around 23.8 hours (Dubowy and Sehgal, 2017).

59 The LNV neurons produce the neuropeptide pigment dispersing factor (PDF) which acts to  
60 synchronise activity throughout the clock circuit (Shafer and Yao, 2014). PDF acts through the PDF  
61 receptor which has broad expression in the circadian network (Im and Taghert, 2010), with rhythmic  
62 synaptic release of the PDF neuropeptide required for maintaining circadian rhythmicity under  
63 constant conditions . The PDF neurons have been termed the ‘morning’ cells due to the absence of  
64 the morning (but not evening) peak of activity in flies either having mutations in the *pdf* gene or  
65 lacking the PDF neurons (Renn et al., 1999). Another group of clock neurons, the LNDs (dorsal lateral  
66 neurons) as well as the PDF-negative 5<sup>th</sup> s-LNvs, have been termed the “evening” cells as they are  
67 necessary for the evening anticipation activity (Grima et al., 2004; Stoleru et al., 2004).

68 In *Drosophila* ageing has been shown to cause reduced and weakened circadian activity in  
69 behaviour, associated with declining levels of PDF (Umezaki et al., 2012). Disruption of the clock has  
70 also been shown to accelerate ageing, in flies lacking a functional clock (Hendricks et al., 2003;  
71 Krishnan et al., 2012; Vaccaro et al., 2017), or keeping flies under mismatched lighting conditions  
72 (Klarsfeld and Rouyer, 1998; Pittendrigh and Minis, 1972; Vaccaro et al., 2016). Studies on how  
73 ageing affects the molecular clock have reported conflicting results: it has been found to remain

74 robust in aged flies (Luo et al., 2012), and to significantly decline in strength with age (Rakshit et al.,  
75 2012). To date no studies have been published on the effect of ageing on the electrical activity of  
76 clock neurons in *Drosophila*. In mice ageing has been shown to result in reduced amplitude of daily  
77 electrical rhythms, measured *in vivo* using multiunit recordings (Nakamura et al., 2011) or from  
78 single cells in slice preparations (Farajnia et al., 2015, 2012).

79 To investigate the effect of ageing on circadian rhythms we took advantage of the *Drosophila* model  
80 that allows systematic monitoring of circadian and sleep behaviour simultaneous from flies across  
81 the range of lifespan. Furthermore, we determine the effect of ageing at the neuronal activity level  
82 by making patch-clamp recordings from the large-LNv clock neurons from young and aged flies.

83

## 84 **2. Materials and methods**

### 85 **2.1. Fly strains**

86 The following fly stocks and their original sources were used, *Pdf::RFP* (Ruben et al., 2012), *iso31*  
87 (Ryder et al., 2004), *Pdf-Gal4* (Bloomington stock centre, #6900) (Park and Hall, 1998) and *UAS-*  
88 *mCD8::GFP* (Bloomington Stock Centre, #5137).

89 All flies were reared on a standard medium based upon the following recipe; 10L batches containing:  
90 400 ml malt extract, 200 ml molasses, 400 g polenta, 90 g active dried yeast, 50 g soya flour and 35 g  
91 granulated agar, with 40 ml of propionic acid (Sigma-Aldrich, #94425) and 100 ml of nipagin (Sigma-  
92 Aldrich, #H5501) added after cooling. Flies for ageing were collected and flipped onto fresh food  
93 every 5 days and maintained in an incubator at 25°C and humidity of 55-65% with a 12:12 LD cycle.

94

### 95 **2.2. Circadian behaviour analysis**

96 Locomotor activity of individual male flies (aged 1, 8, 15, 22, 29, 36, 43 and 49 days old) was  
97 measured using the *Drosophila* Activity Monitoring (DAM) system (DAM2, Trikinetics Inc, USA). Flies  
98 were transferred into DAM tubes after reaching the desired age and were maintained for 5 days  
99 under 12:12 LD conditions, followed by constant darkness. The first 7 days of DD activity was used  
100 for circadian analysis, with period and rhythmicity analysis performed in MATLAB using the  
101 Flytoolbox (Levine et al., 2002).

102

### 103 **2.3. Anticipation index analysis**

104 The morning and evening anticipation indexes were calculated from the activity of flies across the 5  
105 days of LD activity. Morning anticipation was calculated as previously described (Harrisingh et al.,  
106 2007; Zhang and Emery, 2013). Briefly, the average activity was calculated as the ratio of activity  
107 between ZT21.5-24 compared to ZT17-19.5. Evening anticipation was likewise calculated as the  
108 ratio between ZT9.5-12 compared to ZT5-7.5.

109

### 110 **2.4. Sleep analysis**

111 Sleep data were analysed using the Sleep and Circadian Analysis MATLAB Program (S.C.A.M.P.)  
112 (Donelson et al., 2012). Individual raster plots of activity were viewed, and flies that had died before  
113 the end of the experiment were removed from the data. Data were analysed across the 24-hr  
114 period, the 12 hr ‘light phase’, and the 12 hr ‘dark phase’. Sleep is visualised by plotting the mean  
115 amount of sleep in a 30 min bin against the time of day, averaged for the 5 days of the experiment.  
116 From the raw data of sleep amounts and time, a series of measurements of sleep are calculated,  
117 including; ‘total sleep duration’ – sum of all sleep episodes, ‘number of sleep episodes’ – count of all  
118 sleep episodes, ‘mean sleep episode duration’ – average sleep duration (in mins) and ‘sleep latency’  
119 – the time to the first sleep episodes (in mins).

120

121 **2.5. Electrophysiological recording of clock neurons**

122 Whole-cell current clamp recordings were performed as previously described (Buhl et al., 2016;  
123 Chen et al., 2015). For visualisation of the I-LNvs, *Pdf::RFP* flies were used (Ruben et al., 2012), which  
124 is a transgenic fusion of the *Pdf* promoter and *mRFP1* that specifically labels the LNV neurons. Adult  
125 male flies were maintained under a 12:12 LD cycle, and recordings were made at either ZT7-9 (day  
126 condition) or ZT19-21 (night condition), where ZT0 corresponds to lights-on.

127 Firstly, flies were anaesthetised using CO<sub>2</sub>, before decapitation, and the brain removed by acute  
128 dissection with fine forceps in extracellular saline solution containing (in mM): 101 NaCl, 1 CaCl<sub>2</sub>, 4  
129 MgCl<sub>2</sub>, 3 KCl, 5 glucose, 1.25 NaH<sub>2</sub>PO<sub>4</sub>, 20.7 NaHCO<sub>3</sub> at pH 7.2. The photoreceptors, lamina and as  
130 much membrane as possible were removed and whole brains were transferred to a recording  
131 chamber (ALA scientific) filled with extracellular solution and stably held ventral side up using a  
132 custom-built wire harp. Cells were visualised using an Axio Examiner Z1 (Zeiss) using a 63x water  
133 immersion objective, I-LNvs were identified using 555 nm light generated using a Colibri Examiner  
134 light source (Zeiss). All recordings were performed at room temperature (20-22°C) using thick-  
135 walled borosilicated glass electrodes (1B150F-4, World Precision Instruments) ranging in resistance  
136 from 10-16 MΩ filled with intracellular solution containing (in mM): 102 K-gluconate, 0.085 CaCl<sub>2</sub>, 17  
137 NaCl, 0.94 EGTA, 8.5 HEPES, 4 Mg-ATP and 0.5 Na-GTP at pH 7.2. Data were recorded using an Axon  
138 Multiclamp 700B amplifier, digitised with an Axon Digidata 1440 (sample rate 20 kHz, 10kHz Bessel  
139 filter) and recorded using pClamp (10.5: Molecular Devices, CA, USA). Chemicals were acquired from  
140 Sigma (Poole, UK).

141 The liquid junction potential was calculated as 13 mV and subtracted *post-hoc* from all the  
142 membrane voltages. A cell was included in the analysis if the access resistance was less than 50 MΩ.  
143 Membrane potential (MP) and the spontaneous firing rate (SFR) were measured after stabilising for  
144 2–3 minutes. Membrane input resistance ( $R_{in}$ ) was calculated by injecting hyperpolarising current

145 steps from -40 pA to +5 pA in 5 pA steps and measuring the resulting voltage change. Neuronal  
146 excitability was measured by injecting a 500 ms long positive current pulse in 2 pA increments with  
147 increasing amplitude up to +40 pA and manually counting the resulting spikes.

148

## 149 **2.6. Immunohistochemistry and analysis**

150 Flies were briefly anaesthetised using CO<sub>2</sub> and swiftly decapitated and heads immediately placed  
151 into phosphate-buffer saline (PBS) containing 4% PFA (Image-iT™ Fixative Solution, Thermo Fisher  
152 Scientific # R37814) and 0.008% Triton X-100 (Sigma) and fixed for 45 min at room temperature. For  
153 all steps, tubes were covered by foil to protect tissue from light exposure. Fixed heads were quickly  
154 washed twice in 0.5% PBT (PBS with 0.5% Triton X-100), followed by three 20 min washes in PBT,  
155 before being dissected in 0.1% PBT. Brains were blocked in 5% normal goat serum (NGS, Thermo  
156 Fisher Scientific # 50197Z) for 30 min at room temperature. Brains were then incubated with primary  
157 antibodies in 5% NGS, at 4°C for 36 hr on a rotator with tubes upright.

158 Brains were quickly washed twice in PBT, followed by three 20 min washes in PBT, with tubes  
159 upright on a rotator. Brains were then incubated with secondary antibodies in 5% NGS for three  
160 hours at room temperature, and then overnight at 4°C. Brains were rinsed in 0.1 % PBT, followed by  
161 three 20 min washes in PBT, and rinsed twice in PBS. Brains were then aligned on a microscope slide,  
162 with wells created using imaging spacers (SecureSeal™, Grace Bio-Labs #654002), and then mounted  
163 in Vectashield hard set medium (Vector Laboratories). The mounting media was allowed to harden  
164 for 30 min at room temperature, before storage at 4°C. Coverslip edges were sealed with clear  
165 solvent (CoverGrip™, Biotium #23005).

166

167 Table 1 - Antibodies used and sources

Primary Antibodies	Concentration	Source
<b>Mouse monoclonal anti-PDF</b>	1:200	Developmental Studies Hybridoma Bank, #PDF-C7
<b>Rabbit polyclonal anti-GFP</b>	1:1000	Life Technologies # A11122
<b>Secondary Antibodies</b>		
<b>Alexa Fluor Plus 488 Goat anti-mouse</b>	1:1000	Life Technologies # A32723
<b>Alexa Fluor Plus 555 Goat anti-rabbit</b>	1:100	Life Technologies # A32732

168

169 Brains were imaged using a Leica TCS SP8 AOBS confocal laser scanning microscope attached to a Leica  
 170 DMi8 inverted epifluorescence microscope, equipped with ‘hybrid’ Gallium arsenide phosphide (GaAsP)  
 171 detectors with the green channel imaged at 480 – 551 nm and the red at 571 – 650 nm. We used a 20x  
 172 glycerol immersion objective (HC PL APO CS2, Leica) and obtained confocal stacks with a 2  $\mu$ m step size  
 173 and 512 x 512 pixels. The obtained confocal stacks were analysed using the FIJI implementation of ImageJ  
 174 (Schindelin et al., 2012). Besides contrast, brightness, colour scheme and orientation adjustments, no  
 175 further manipulations were made to the images.

176 To quantify the axonal arbour of the dorsal projections we used an adaptation of the Sholl method (Sholl,  
 177 1953), as has been previously reported (Fernández et al., 2008). Briefly, using six evenly spaced (10  $\mu$ m)  
 178 concentric rings centred at the first branching of the dorsal projections, and counting the number of  
 179 intersections of each projection with the rings. Scoring was performed blind to the experimental  
 180 condition.

181

## 182 **2.7. Statistical analysis**

183 All statistical analyses were performed using Graphpad Prism 7 (Graphpad Software, USA), with an  $\alpha$   
 184 level of  $p < 0.05$  considered significant. Data for ageing experiments showed non-normal distribution  
 185 and were plotted as the median with interquartile range, the non-parametric alternative Kruskal-  
 186 Wallis test was used, with post-hoc analysis conducted using Dunn’s test. For ageing data, statistical  
 187 comparisons between groups were compared to the D1 group.

188 For electrophysiological and imaging experiments, groups were compared using two-way ANOVA  
189 with Tukey's multiple comparisons test, with 'age' and 'time of day' as factors.

190

191 **3. Results**

192 **3.1. Ageing caused a weakening in circadian behavioural output and lengthening of the free-  
193 running period**

194 To address the impact of ageing on the circadian clock we used *Drosophila* to conduct a  
195 comprehensive behavioural analysis of circadian activity of male flies at 1-week intervals during the  
196 ageing process. Flies were first kept for 5 days in a 12hr:12hr light-dark (LD) cycle and showed  
197 normal diurnal behaviour, with morning and evening peaks of activity (Figure 1A). Flies were then  
198 maintained in constant darkness and showed typical free-running behaviour. In agreement with  
199 previously reported work (Rakshit et al., 2012; Umezaki et al., 2012) we found that ageing resulted in  
200 a significant decline in the strength of circadian locomotor activity under free-running conditions  
201 (Figure 1B,  $p=0.0001$ , Kruskal-Wallis statistic = 27.17), with a steep decline in flies aged 36 days and  
202 older, and a significant lengthening of the period of the observed behavioural activity (Figure 1C,  
203  $p<0.0001$ , Kruskal-Wallis statistic = 95.3). We also found there was a significant age-related  
204 reduction in total locomotor activity (Figure 1D,  $p<0.0001$ , Kruskal-Wallis statistic = 37.89).

205

206 **3.2. Ageing alters daily activity structure in light-dark conditions and reduces anticipatory  
207 behaviour**

208 Given that there was a reduction in the amount of locomotor activity in older flies (Figure 1D), we  
209 sought to further examine how the daily structure of activity under normal LD conditions was altered  
210 by the ageing process. Some of the hallmark features of daily activity of male flies recorded using the  
211 *Drosophila* Activity Monitoring (DAM) system are the morning and evening peaks in locomotor

212 activity (Dubowy and Sehgal, 2017) and anticipation of the light-dark transition (see Figure 2A). Both  
213 the peaks of activity and anticipation behaviour are affected by manipulations of the circadian  
214 system (Lear et al., 2009). We investigated the effect of age on morning and evening anticipation by  
215 first normalising locomotor activity for an individual fly to be the percentage of the daily total (Figure  
216 2B). The anticipation index was then quantified as the proportion of an individual fly's daily activity  
217 occurring in the 2.5 hrs immediately prior to the light-dark transition compared to the 2.5 hrs in the  
218 middle of the day/night (Harris et al., 2007). Older flies showed significant reductions morning  
219 anticipation index compared to young flies ( $p<0.0001$ , Kruskal-Wallis statistic = 28.93), and a slight  
220 reduction in the evening anticipation index ( $p<0.0001$ , Kruskal-Wallis statistic = 32.45) (Figure 2C).

221

### 222 **3.3. Ageing alters the daily structure of sleep**

223 Given that there is a strong connection between the circadian clock and sleep, we sought to also  
224 investigate how sleep is altered by age. Sleep analysis was performed for the five days under a LD  
225 cycle at the start of the circadian experiment, with sleep classified under the common convention of  
226 periods of immobility longer than five minutes in duration (Hendricks et al., 2000; Shaw et al., 2000).  
  
227 The daily structure of sleep in older male flies was noticeably different to that of young flies (Figure  
228 3A) with a visible increase in the amount of daytime sleep and a shift towards sleep earlier in the  
229 day. Quantification of total sleep showed a significant effect of age ( $p=0.0006$ , Kruskal-Wallis statistic  
230 = 25.7) (Figure 3B). Looking only at sleep during the daytime (Figure 3C) there was a significant  
231 increase with age ( $p<0.0001$ , Kruskal-Wallis statistic = 33.83), however there was no effect of age on  
232 night-time sleep ( $p=0.31$  Kruskal-Wallis statistic = 8.253) (Figure 3D). Measuring the latency of sleep  
233 after the LD transitions demonstrated a significant reduction in the speed at which older flies started  
234 sleeping both during the day ( $p<0.0001$ , Kruskal-Wallis statistic = 102.7) (Figure 3E) and night  
235 ( $p<0.0001$ , Kruskal-Wallis statistic = 61.33) (Figure 3F). Analysing the parameters of sleep episodes,  
236 we found that there was a significant increase in the number of sleep episodes with age ( $p<0.0001$ ,

237 Kruskal-Wallis statistic = 58.79) (Figure 3G) and a significant difference in mean sleep episode  
238 duration ( $p < 0.0001$ , Kruskal-Wallis statistic = 32.56), (Figure 3H).

239

240 **3.4. Electrical properties of clock neurons are altered by ageing**

241 We have demonstrated that the ageing process causes significant changes to the behavioural  
242 outputs of the circadian clock circuit of *Drosophila* and therefore set out to investigate if these were  
243 underpinned by changes in clock neuronal activity. To measure the effects of ageing on clock  
244 neurons we made recordings from the prominent wake-promoting I-LNv arousal, the most  
245 accessible and well-studied group of clock neurons in *Drosophila* (Buhl et al., 2016; Cao and  
246 Nitabach, 2008; Parisky et al., 2008; Sheeba et al., 2008). Recordings were made during the day and  
247 at night in explant brain preparations made from young (day (d) 1-5) and middle-aged (d28-35) flies  
248 (Figure 4) and measured the electrophysiological properties of these cells (Figure 5). Recordings  
249 from flies older than 35 days were limited due to the technical difficulties making stable recordings  
250 from aged neurons, with older brains being more difficult to dissect cleanly and difficulties to  
251 achieve good seals and access due to changes in the older membranes.

252

253 As previously reported, young I-LNvs showed a strong day-night difference in both their spontaneous  
254 firing rate (SFR) and membrane potential (MP) (Cao and Nitabach, 2008; Chen et al., 2015; Sheeba et  
255 al., 2007), but the response to an injected current pulse or the input resistance did not differ  
256 significantly between day and night (Buhl et al., 2016) (Figure 5). Here we report that the diurnal  
257 modulation of SFR and MP are maintained in the I-LNvs recorded from 28-35 day old flies, with no  
258 difference found in the response to a current injection between young and aged flies. Interestingly  
259 we report a significant decrease in the input resistance of aged I-LNvs (Figure 5A), indicative of an  
260 increase in overall conductance across the membrane.

261

262 **3.5. s-LNv terminal remodelling is reduced by ageing**

263 It has previously been demonstrated that the dorsal projections of the s-LNv neurons show daily  
264 remodelling in complexity under clock control (Fernández et al., 2008). To test if this was still  
265 occurring in older flies, day/night changes in PDF terminal morphology were measured in flies aged  
266 30 days (Figure 6). Using the previously published protocol (Fernández et al., 2008) we found that  
267 the remodelling was no longer a significant feature in aged brains. There was a significant overall  
268 effect of time of day ( $p=0.0045$ , two-way ANOVA,  $F(1,24)=9.822$ ) but no effect of age ( $p=0.5286$ ,  
269 two-way ANOVA,  $F(1,24)=0.4088$ ). Multiple comparisons tests showed the magnitude of the day-  
270 night difference was reduced in older flies and no longer being significantly different between day  
271 and night ( $p=0.6138$ , Tukey's test,  $DF=24$ ,  $q=1.741$ ) (Figure 6C).

272

273 **4. Discussion**

274 Ageing is known to have a significant impact on circadian behaviour but what effect this has at a  
275 neuronal level is poorly understood. In this study we have conducted a systematic analysis of the  
276 effect of ageing on circadian behaviour and related this to electrical activity of I-LNv clock neurons  
277 finding a significant reduction in the input resistance of aged neurons.

278 Our behavioural experiments complement the work of previous studies in showing that the strength  
279 of the free-running behaviour weakens and period lengthens with age (Umezaki et al., 2012). In  
280 addition, we go further by using a systematic approach to monitor flies at 1-week intervals across  
281 the ageing process and show that there is an age-dependent decrease in rhythm strength (Figure 1B)  
282 and an equivalent increase in period length with age (Figure 1C). Mouse experiments have found  
283 that ageing results in a lengthening in period in both behavioural activity (Turek et al., 1995;  
284 Valentinuzzi et al., 1997) and in molecular rhythms in the SCN (Nakamura et al., 2015).

285 We further sought to investigate how the daily structure of activity under light-dark conditions is  
286 altered by ageing, by quantifying changes in morning and evening anticipatory activity (Figure 2). We

287 found that there was a significant effect of age on both the morning and evening anticipation  
288 indexes (Figure 2C), with a greater reduction in the morning peak. L<sub>Nv</sub> neurons are required for  
289 correct morning anticipation (Grima et al., 2004; Stoleru et al., 2004) and are obvious candidates for  
290 involvement in an age related decline in this anticipatory behaviour. Morning anticipatory behaviour  
291 is also linked to expression of PDF, with *pdf*<sup>01</sup> and *PDF-RNAi* flies showing significant reductions in  
292 morning anticipation (Shafer and Taghert, 2009). A reduction in PDF expression in aged flies has  
293 previously been demonstrated (Umezaki et al., 2012), providing further evidence for the importance  
294 of PDF in maintaining healthy rhythms with age, and supporting a hypothesis that reduced PDF  
295 signalling with age underlies the weakening of behavioural rhythmicity.

296 The I-L<sub>Nv</sub> neurons are involved in promoting arousal (Chung et al., 2009; Sheeba et al., 2008) and  
297 regulating sleep and latency during the early night (Liu et al., 2014). We made use of the DAM  
298 recording system to monitor sleep under light-dark conditions, using the widely accepted definition  
299 of sleep as period of immobility greater than 5 mins (Shaw et al., 2000). Ageing is known to cause  
300 changes in the sleep profile across many organisms including mice (Valentinuzzi et al., 1997), non-  
301 human primates (Zhdanova et al., 2011) and humans (Moraes et al., 2014). Previous *Drosophila*  
302 studies on the effects of ageing on sleep have reported that sleep becomes more fragmented with  
303 age (Koh et al., 2006; Vienne et al., 2016), showing a similar increase in sleep episode number and  
304 decrease in mean sleep episode duration compared to our results (Figure 3G & 3H).

305 Electrical silencing of L<sub>Nv</sub> neurons causes deficits in free-running clock behaviour (Depetris-Chauvin  
306 et al., 2011), demonstrating a link between electrical activity and behaviour. Most  
307 electrophysiological studies use young flies aged between 3 and 7 days for recordings (Cao and  
308 Nitabach, 2008), with a limited amount of recordings made from 25 day old flies only looking at the  
309 active firing properties of the neurons (Sheeba et al., 2007). Here we report the effect of ageing on I-  
310 L<sub>Nv</sub> neuronal activity and electrical properties. We perform whole-cell patch clamp recordings from  
311 young and aged neurons and report no major differences in the observed spontaneous activity of I-

312 LNV neurons (Figure 4). Further analysis of the electrical properties of I-LNV neurons showed that  
313 there was a significant effect of age in reducing the input resistance, which surprisingly did not affect  
314 spontaneous firing rate, membrane potential or excitability (Figure 5). We propose the age-related  
315 changes in I-LNV properties are linked with the observed changes in activity and sleep during light-  
316 dark conditions.

317 There are multiple possible explanations for a decrease in input resistance without changing the  
318 active properties of the neurons. One hypothesis would be the involvement of chloride ( $\text{Cl}^-$ )  
319 channels, which could become open and decrease resistance without changing the membrane  
320 potential, alternatively, the observed reduction in input resistance could result from age-related  
321 changes in the composition of ion channels in the membrane, with future experiments needed to  
322 evaluate between potential hypotheses. The I-LNV express the GABA<sub>A</sub> receptor *Resistant to dieldrin*  
323 (*Rdl*), which when activated by GABA selectively conducts  $\text{Cl}^-$  through its pore. *Rdl* has important  
324 roles in promoting sleep, with a mutation in *Rdl* that causes extended channel openings resulting in  
325 increased sleep duration and decreased latency (Agosto et al., 2008; Parisky et al., 2008).  
326 Conversely, knocking down the *Rdl* gene in the PDF neurons reduces sleep, again suggesting GABA  
327 regulates sleep through the LNvs and *Rdl* receptor function (Chung et al., 2009). Therefore, it is  
328 possible that during ageing there is an increase in GABA activation through *Rdl* in the I-LNvs, causing  
329 increased  $\text{Cl}^-$  conductance. This increase in  $\text{Cl}^-$  conductance may contribute to the observed  
330 reduction in input resistance recorded and also drive the increase in sleep duration and decreased  
331 sleep latency in aged flies.

332 Studies of ageing on electrical activity of mouse clock neurons found no effect of age on input  
333 resistance but reveal a reduction in the difference between day and night firing rates (Farajnia et al.,  
334 2012), showing differences of the effects of ageing between different clock neurons in *Drosophila*  
335 and mouse.

336 Our electrophysiological experiments were limited to the I-LNvs so we can only link the changes in  
337 neuronal properties we observed to the changes in morning activity and sleep in LD conditions as  
338 the I-LNv do not maintain molecular oscillations in DD (Grima et al., 2004), although it is possible  
339 that similar changes in neuronal properties are occurring in the s-LNvs where molecular oscillations  
340 do persist in constant conditions. We sought to investigate changes to the s-LNv neurons, namely  
341 the remodelling of the s-LNv dorsal projections. Analysis of the branching of the s-LNv projections  
342 demonstrated that the day-night difference in complexity is reduced by ageing (Figure 6), indicating  
343 changes in the distribution of the PDF release network in older flies. Given the role of PDF in  
344 regulating the activity of different groups of clock neurons, namely through excitation of dorsal clock  
345 neurons (Seluzicki et al., 2014), changes in PDF signalling would contribute to changes in the clock  
346 network as a whole. The s-LNv neurons are known to be important for maintaining behavioural  
347 rhythmicity under constant conditions and we propose this weakening of s-LNv terminal remodelling  
348 underlies the age-related weakening in circadian locomotor behaviour.

349 Our study builds upon the existing literature demonstrating an age-dependent decline in circadian  
350 behavioural outputs, and importantly links this to changes in the electrophysiological and structural  
351 properties of clock neurons. Further work is necessary to fully understand what the implications of  
352 these changes are for the circadian clock network as a whole and if similar changes are occurring in  
353 other groups of clock neurons in *Drosophila*.

354

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552 Figure 1. Locomotor activity at different timepoints in the ageing process. (A) Top panel - group  
553 activity profiles during LD from wild type flies. Middle-panel - double plotted actograms of  
554 representative individual flies are shown, activity scaled to individual maximum. Flies were  
555 maintained for 5 days of LD before being maintained in constant darkness (DD). In the actograms  
556 white represents day, grey background represents darkness. Bottom panel – group activity profile  
557 during the 5<sup>th</sup> day of DD (DD5). (B) Circadian behaviour weakens with age as measured by the  
558 rhythmicity statistic. (C) Ageing causes lengthening of period in wild type flies. (D) Average daily  
559 locomotor activity during LD is significantly reduced by ageing. On the x axis D signifies days after  
560 eclosion. ‘\*\*\*’ represents p<0.001 as determined using the Kruskal-Wallis test with Dunn’s post hoc  
561 test, data plotted as median with error bars representing the interquartile range.

562  
563 Figure 2. Morning and evening anticipation reduce with age in wild type flies. (A) Schematic of  
564 morning and evening anticipation index. (B) Normalised daily group activity plots of 1, 15, 29 and 43-  
565 day old flies. (C) Quantification of anticipation index shows that the morning anticipation index is  
566 significantly reduced by age, and that evening anticipation is slightly reduced. N=20-32 for each  
567 group, ‘\*’ represents p<0.05, ‘\*\*’ - p<0.01, ‘\*\*\*’ - p<0.001 as determined using the Kruskal-Wallis  
568 test with Dunn’s post hoc test, data plotted as median with error bars representing the interquartile  
569 range.

570  
571 Figure 3. Ageing alters the daily structure of sleep in *Drosophila*. (A) Daily sleep profile of groups of  
572 male flies aged 1, 29 and 43 days old, average across 5 days. Shaded area represents the 95%  
573 confidence interval. (B – H) Quantification of sleep parameters for flies aged D1, 8 ,15, 22, 29, 36, 43  
574 and 49 days, flies were monitored in parallel under identical conditions. Error bars plot the median  
575 and interquartile range with ‘\*’ representing p<0.05, ‘\*\*’ - p<0.01, ‘\*\*\*’ - p<0.001 with statistical  
576 testing performed by Kruskal-Wallis test with Dunn’s multiple comparisons. (B) Mean total sleep  
577 duration (C) Mean daytime sleep (D) Mean night-time sleep (E) Latency to sleep after lights on (F)

578 Latency to sleep after lights off (G) Number of sleep episodes (H) Sleep episode duration (see  
579 methods for definitions).

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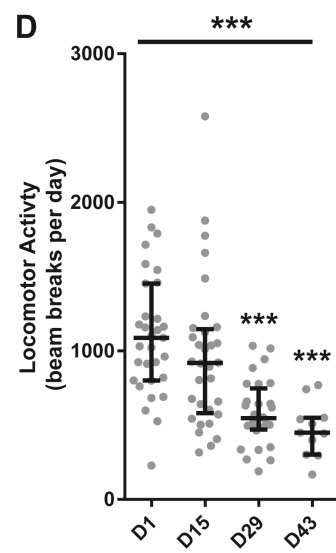
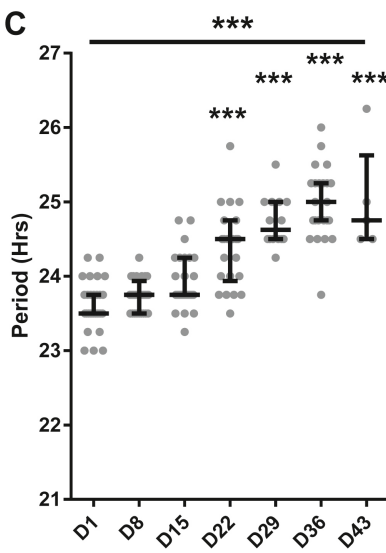
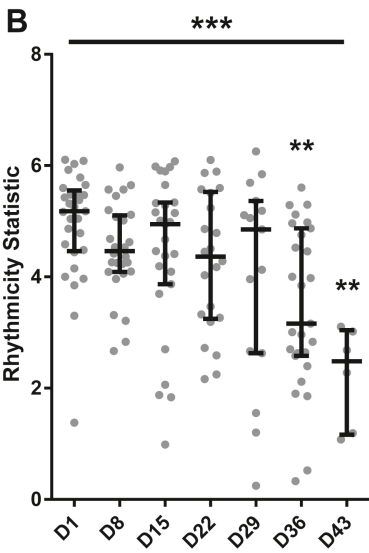
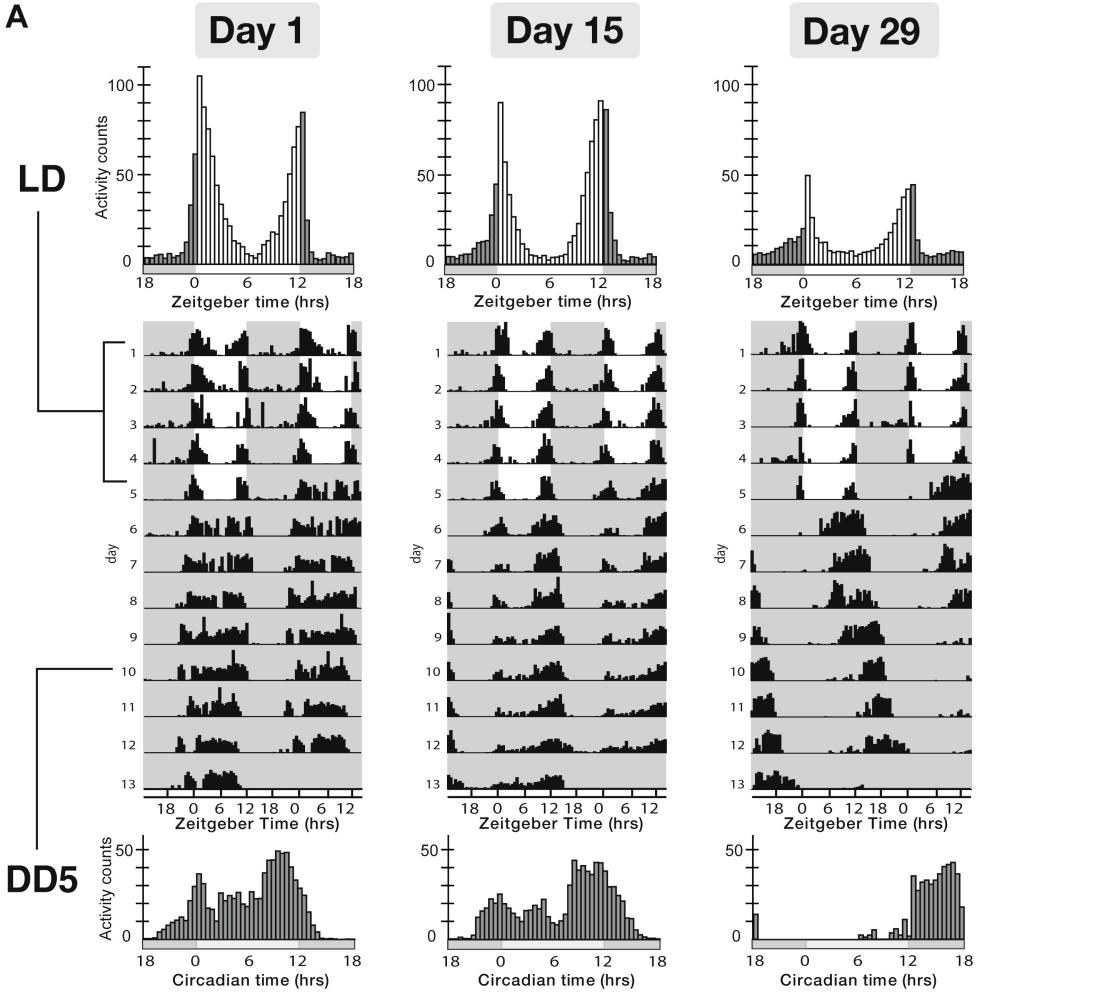
581 Figure 4 Electrophysiological characterisation of I-LNv clock neurons. Membrane potential and  
582 spontaneous activity (left panels) and firing response to a current pulse (right panels, colour-coded  
583 as indicated) of wild type I-LNvs from young (d1-5) and aged (d28-35) flies recorded at day (ZT 7-9)  
584 and night (ZT 19-21).

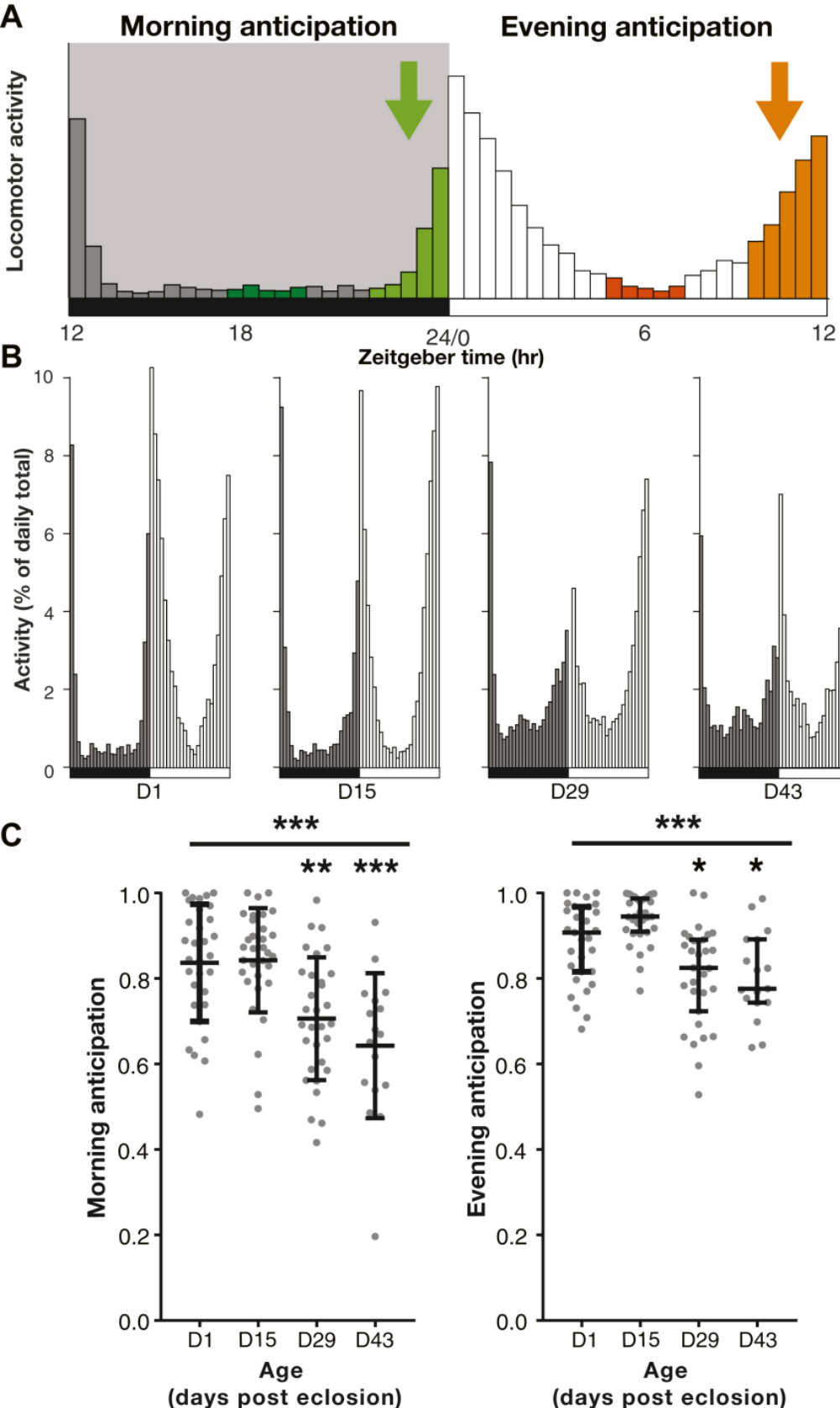
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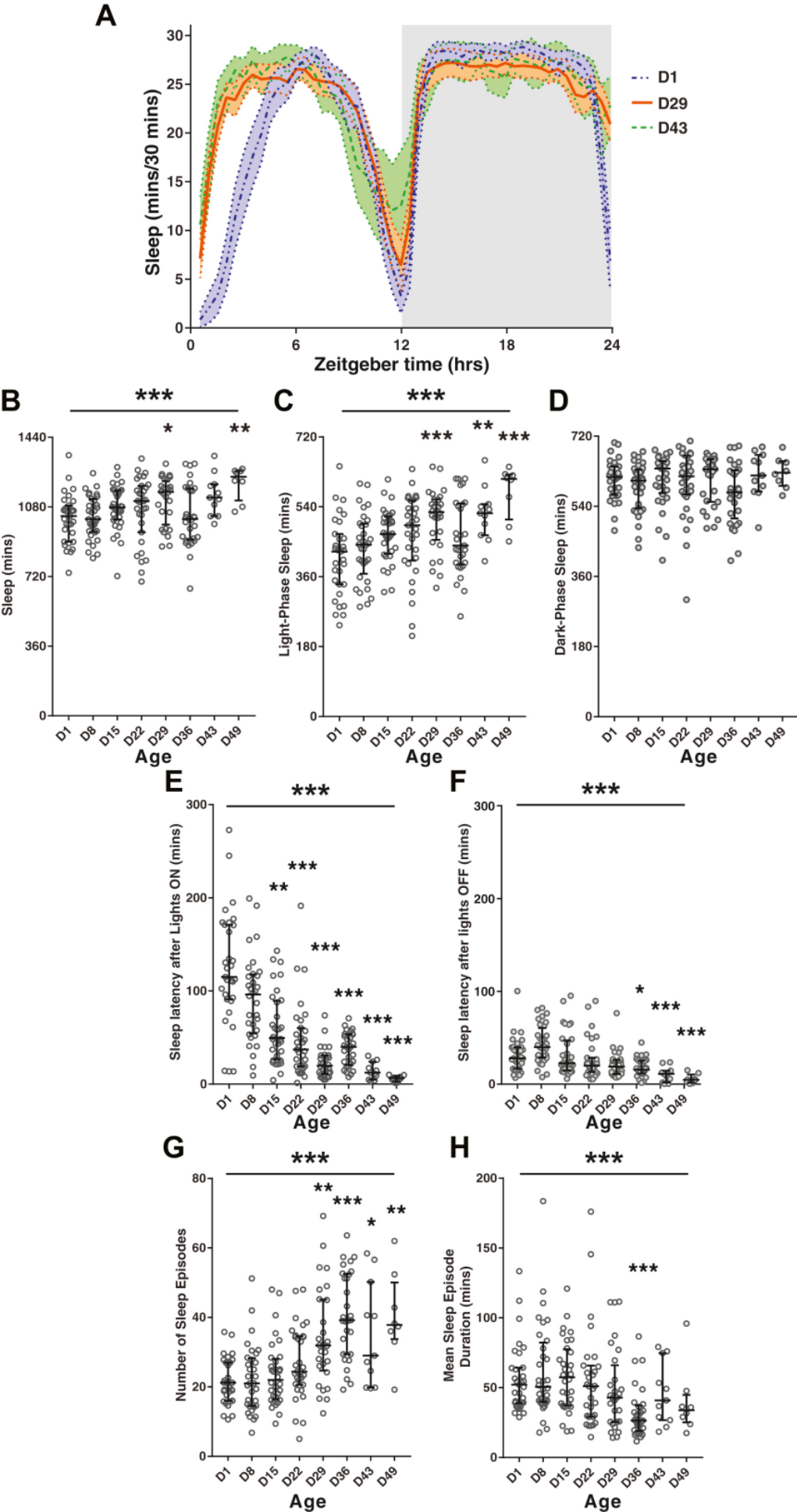
586 Figure 5. Quantitative analysis of electrophysiological properties of I-LNv clock neurons from young  
587 (d1-5) and middle-aged (d28-35) flies in day and night conditions. (A) Analysis of input resistance  
588 ( $R_{in}$ ) showed a highly significant effect of age ( $p<0.0001$ ) and an effect of time of day ( $p<0.005$ ) but  
589 no interaction. (B) Analysis of the spontaneous firing rate (SFR) showed a significant effect of time of  
590 day ( $p <0.0001$ ) but no effect of age. (C) Analysis of the membrane potential (MP) values showed a  
591 significant effect of time of day ( $p<0.0001$ ). (D) Analysis of the responses to an injected current pulse  
592 ( $f_{+40pA}$ ) showed no significant effects. Data were analysed using two-way ANOVA with Tukey's  
593 multiple comparisons test, error bars show the mean  $\pm$  SEM. Each data point represents a single I-  
594 LNV neuron.

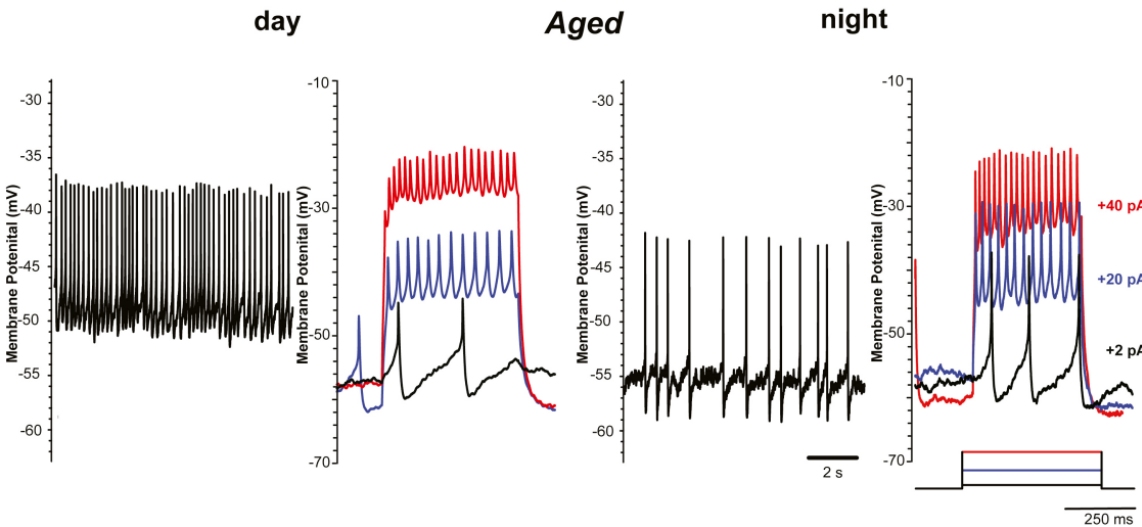
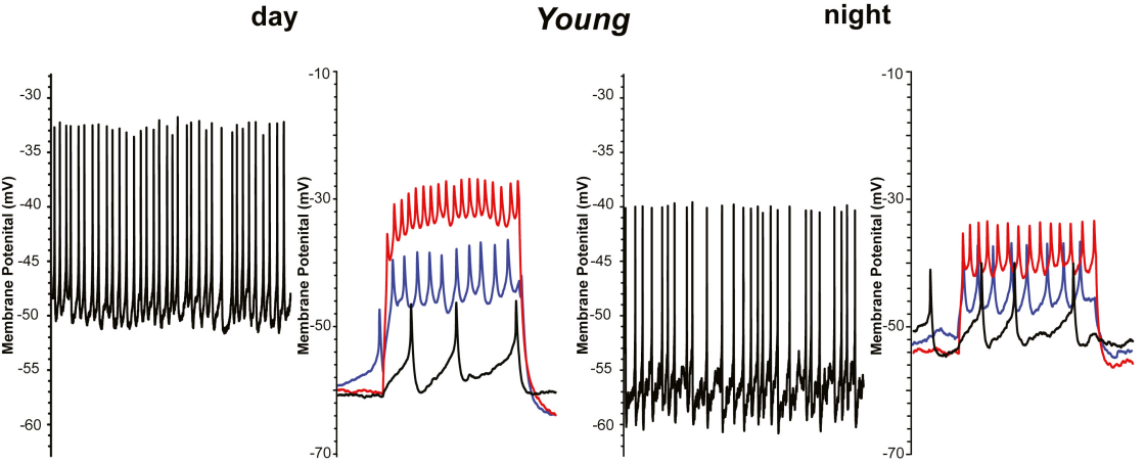
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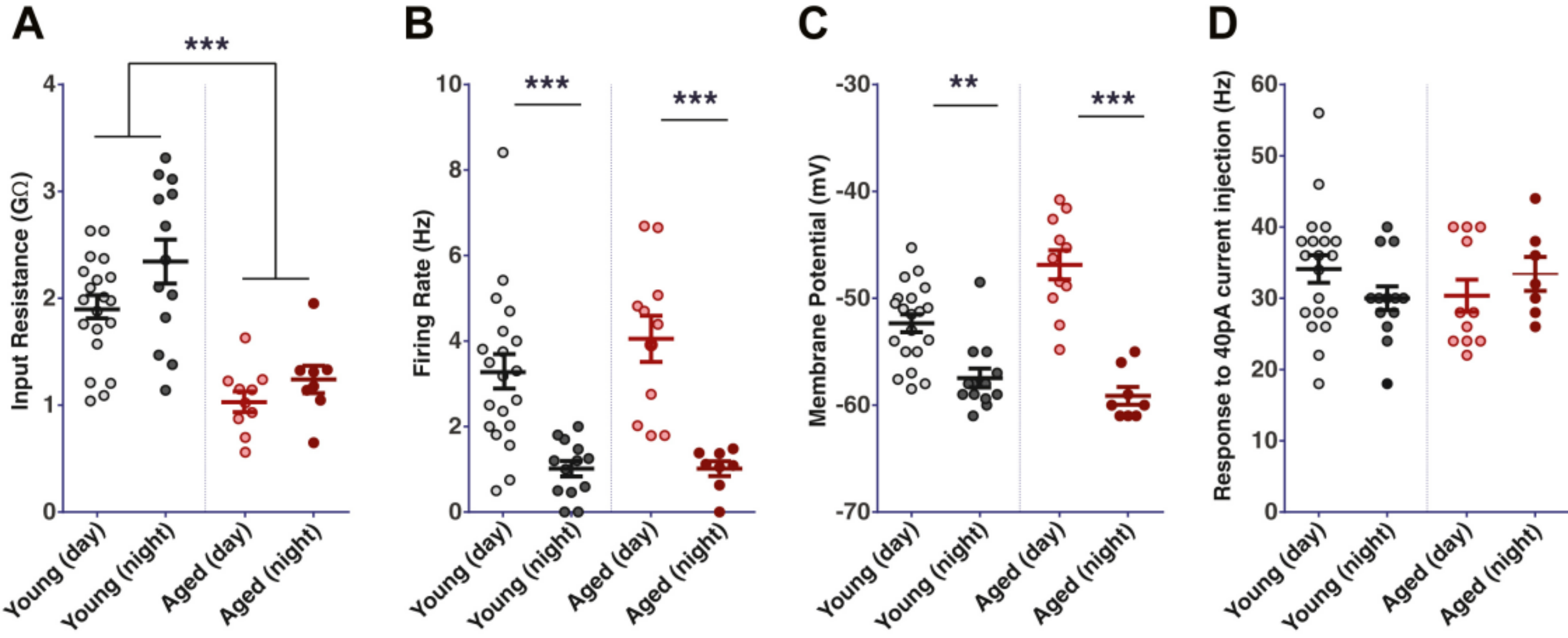
596 Figure 6. Daily reorganization in the PDF terminals is reduced by ageing. (A) *pdf>mCD8-GFP* wild type  
597 brains dissected at ZT2 and ZT14. Brains were stained with anti-GFP (green) and anti-PDF (magenta)  
598 antibodies. Scale bar = 50  $\mu$ m. (B) Schematic depiction of how the quantification of the PDF axonal  
599 branching was carried out. (C) The total number of intersections between the concentric rings and  
600 the axonal projections was quantified and showed daily remodelling. Error bars show mean  $\pm$  SEM,  
601 statistical analysis performed by two-way ANOVA with Tukey's multiple comparisons test. N>6 for all  
602 groups, quantification was performed on the dorsal projections originating from a group of s-LNV  
603 neurons.

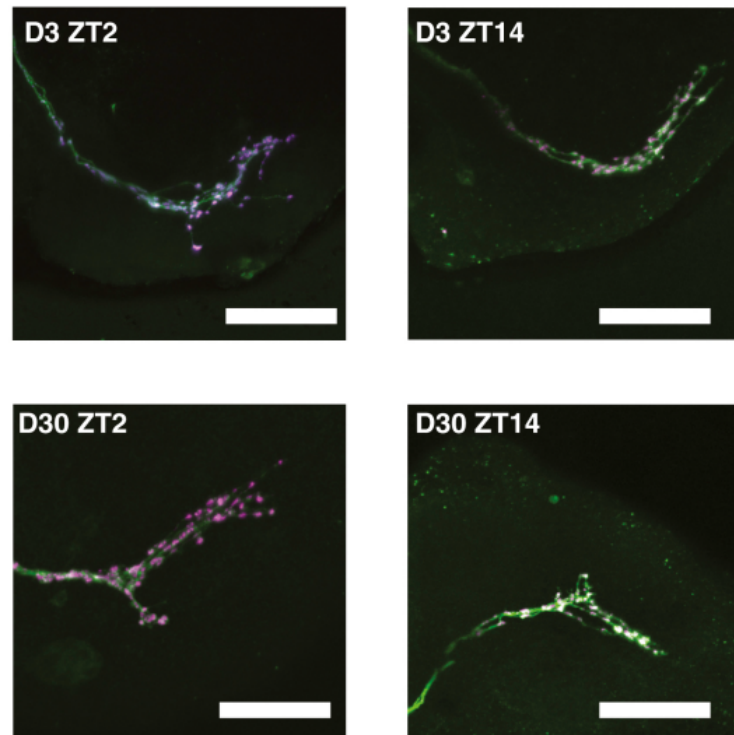
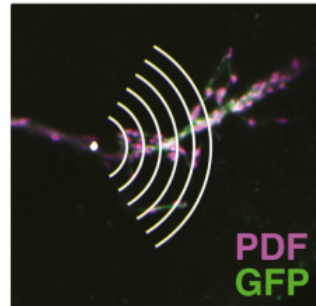










**A****B****C**