

脳梁軸索投射ニューロンは特異的な資格応答と局所結合様式を示す

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1 **Title: Long-range interhemispheric projection neurons show biased response**
2 **properties and fine-scale local subnetworks in mouse visual cortex**
3 **Running title: Response and connectivity profiles of visual callosal neurons**

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31 **Abstract**

32 Integration of information processed separately in distributed brain regions is essential for
33 brain functions. This integration is enabled by long-range projection neurons, and further,
34 concerted interactions between long-range projections and local microcircuits are crucial.
35 It is not well known, however, how this interaction is implemented in cortical circuits. Here,
36 to decipher this logic, using callosal projection neurons (CPNs) in layer 2/3 of the mouse
37 visual cortex as a model of long-range projections, we found that CPNs exhibited distinct
38 response properties and fine-scale local connectivity patterns. *In vivo* 2-photon calcium
39 imaging revealed that CPNs showed a higher ipsilateral (to their somata) eye preference,
40 and that CPN pairs showed stronger signal/noise correlation than random pairs. Slice
41 recordings showed CPNs were preferentially connected to CPNs, demonstrating the
42 existence of projection target-dependent fine-scale subnetworks. Collectively, our results
43 suggest that long-range projection target predicts response properties and local
44 connectivity of cortical projection neurons.

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46 **Keywords: Callosal projection neuron; visual cortex; ocular dominance; 2-photon**
47 **imaging; slice physiology**

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53 **Introduction**

54 The cerebral cortex processes information arriving from the external world in a distributed

55 manner. To achieve this, there must be biophysical substrates enabling proper routing, local
56 computation, and integration of information. Integration of distributed information is
57 mediated by cortical projection neurons; thus, the functional organization of these neurons
58 have been extensively studied (Glickfeld et al., 2013; Kim et al., 2015; Lur et al., 2016;
59 Movshon and Newsome, 1996; Sato and Svoboda, 2010; Yamashita et al., 2013). Such studies
60 have revealed that as a general principle, neurons projecting to different areas encode
61 different features of the sensory input. In addition, fine-scale local network motifs (Brown
62 and Hestrin, 2009; Kampa et al., 2006; Otsuka and Kawaguchi, 2011; Yoshimura et al., 2005)
63 implicated in selective local computation have been identified. Recent work has found that
64 cortical neurons that do not share their projection targets are rarely connected to each other,
65 suggesting the existence of strongly segregated local network motifs suited for highly
66 selective information transmission (Kim et al., 2018). However, the organizational principles
67 linking projection targets to functional properties and fine-scale local networks have not
68 been fully addressed. More specifically, whether the strongly segregated local network is
69 the general motif for all the long-range projection populations is an open question to be
70 addressed.

71 Callosal projection neurons (CPNs) are essential for integrating information
72 processed in separate hemispheres (Hubel and Wiesel, 1967; Van Essen et al., 1982), and
73 thus serve as an ideal model to study the functional organization of long-range projection
74 neurons. Although many lesion and inactivation studies have been performed, to what
75 extent callosal inputs influence the visual response properties of primary visual cortex (V1)
76 neurons is still controversial (Berlucchi and Rizzolatti, 1968; Cerri et al., 2010; Dehmelt and
77 Lowel, 2014; Minciacchi and Antonini, 1984; Payne et al., 1980; Ramachandra et al., 2020;

78 Restani et al., 2009; Schmidt et al., 2010; Wunderle et al., 2013; Zhao et al., 2013). Due to
79 the lack of a method enabling selective *in vivo* recordings from CPNs and non-CPNs, the
80 specific visual information encoded and transmitted by individual CPNs remains largely
81 unknown. Considering the versatility of CPNs as experimental models such as for cortical
82 area identification (Wang and Burkhalter, 2007), for optogenetic circuit mapping *in vitro*
83 (Petreanu et al., 2007), for antidromic optogenetic stimulation *in vivo* (Sato et al., 2014), for
84 studying bilaterally concerted behavior such as eye-movement (Itokazu et al., 2018; Sato
85 et al., 2019), and for developmental circuit formation (Mizuno et al., 2007), functional
86 characterization of CPNs itself is also of high interest.

87 In higher mammals such as cats and monkeys, the inputs from each eye are
88 spatially segregated in V1 as ocular dominance columns (Van Essen et al., 1992; Wiesel and
89 Hubel, 1965). On the other hand, rodents, especially mice have no ocular dominance
90 columns ((Antonini et al., 1999; Drager, 1974; Gordon and Stryker, 1996; Mrsic-Flogel et al.,
91 2007; Scholl et al., 2015) but see (Laing et al., 2015) for results in the Long Evans rat) rather,
92 functionally distinct neuronal populations are spatially organized in a “salt and pepper”
93 manner (Ohki and Reid, 2007). Thus, to assess the relationships between CPN response
94 properties and local connection patterns, it is crucial to assess neuronal responses and local
95 connectivity at single-cell resolution and to directly compare the functional organization of
96 CPNs to surrounding non-CPNs. Here, we achieve this by combining retrograde-labeling of
97 CPNs with *in vivo* two-photon calcium imaging and *ex vivo* slice recordings in layer 2/3 (L2/3)
98 of the mouse visual cortex.

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119 **Materials and Methods**

120 **LEAD CONTACT AND MATERIALS AVAILABILITY**

121 Further information and requests for resources and reagents should be directed to and will
122 be fulfilled by the Lead Contact, Kenta M. Hagihara (hagi_k@med.kyushu-u.ac.jp; kenta-
123 hagihara@umin.ac.jp).

124

125 **EXPERIMENTAL MODEL AND SUBJECT DETAILS**

126 All of the experiments were performed in accordance with the institutional animal welfare
127 guidelines of the Animal Care and Use Committee of Kyushu University, Kyoto University
128 and National Institute for Physiological Sciences, and they were approved by the Ethical
129 Committee of Kyushu University, the local committee for handling experimental animals in
130 the Graduate School of Science, Kyoto University, and National Institute for Physiological
131 Sciences.

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133 **METHOD DETAILS**

134 *Stereotactic retrograde tracer injection for imaging experiments and histology*

135 Mice (C57BL/6, both males and females, aged at postnatal day 60-90) were housed in the
136 temperature-controlled animal room with 12h/12h light/dark cycle. Cholera Toxin Subunit
137 B Alexa Fluor 555 or 488 Conjugate (CTB555 or CTB488, 1.0%, weight/volume, Invitrogen)
138 and Green Retrobeads™ IX (Green Beads, Lumafuor Inc.) were used as retrograde tracers
139 (Conte et al., 2009; Li et al., 2015). The mice were subjected to stereotactic tracer injections
140 using pulled glass micropipettes. The mice were anesthetized with 440 mg/kg chloral
141 hydrate (Tokyo Chemical Industry) by intraperitoneal injection. The mice were then fixed to
142 a small-animal stereotactic device (Narishige). The head skin was cut at the midline, and the
143 periosteum was removed using a surgical knife. The skull was thinned with a drill, and a
144 small craniotomy was made using a 30- gauge needle. The tracer was injected through a
145 pulled glass micropipette connected to a Hamilton syringe (Hamilton Company), which was
146 pumped using a syringe pump device (World Precision Instruments). The stereotactic

147 injections were administered at BZ. The stereotaxic coordinates for BZ injection were 3.5
148 mm lateral of the midline and 2.0 mm in front of the anterior margin of the transverse sinus.
149 The tracer solution was injected at a rate of 0.05–0.1 $\mu\text{l}/\text{min}$ at a volume of 2.0 μl to cover
150 most of BZ (**Figure S1A**), and after the injection, the pipette was held in place for an
151 additional 10 min before removal. After the removal of the micropipette, the skin incision
152 was sutured. The post-injection animals were housed normally for 5–14 days before
153 imaging or histology experiments. Fluorescent histological images were acquired using a
154 confocal laser-scanning microscope (LSM700, Zeiss).

155 In the experiment described in **Figure S2**, a cocktail of Green Beads and CTB555
156 (1.0%) was used. Given the efficiency (0-1) of Green Beads and of CTB555 as a and b ,
157 respectively, and the fraction of CPNs as n , the fraction of CPNs labeled with both Green
158 Beads and CTB555 would be $n*a*b$. Considering that neurons labeled with Green Beads
159 were also mostly labeled with CTB555, here $n*a*b$ can be regarded as near $n*a$ and thus b
160 can be regarded as 1. Thus, in this system, the efficiency of CTB555 can be regarded as
161 almost 100% and neurons not labeled with CTB555 can be regarded as non-CPNs.

162

163 *Stereotactic retrograde tracer injection for slice experiments*

164 For retrograde labeling of CPNs, 28–32-day old C57BL/6 mice of either sex were
165 anesthetized with propofol (0.2%, 100 ml/kg ip, Maruishi) and then fixed to a small-animal
166 stereotactic device. CTB488 or CTB555 (300–500 nl) was pressure-injected (Toohey Spritzer
167 microinjector, Toohey Company) into the visual cortex contralateral to the eventual
168 recording site (3.5 mm lateral to the midline, 1.5 mm rostral to the anterior margin of the
169 transverse sinus, 300–600 μm in depth) with pulled glass pipettes (tip diameter 10–20 μm ,

170 Narishige) to cover most of BZ. The pipette was held in place for 10 min before and after
171 injection. After removal of the pipette, the skin incision was sutured and mice were allowed
172 to recover for 4–8 days before recording.

173

174 *Animal preparation and surgery for in vivo calcium imaging*

175 5-14 days after CTB555 injection, the animals were prepared for *in vivo* calcium imaging as
176 previously described (Hagihara et al., 2015; Ohki and Reid, 2014). In brief, mice were
177 anesthetized with isoflurane, then a custom-made metal plate was mounted on the skull,
178 and a craniotomy was carefully performed on the other hemisphere above the area where
179 fluorescence from CTB555 was clearly observed through the skull. We dissolved 0.8 mM
180 Oregon Green 488 BAPTA-1 AM (OGB1) in DMSO with 20% pluronic acid and mixed it with
181 ACSF containing 0.05 mM Alexa594 (all obtained from Invitrogen). A glass pipette (3–5 μm
182 tip diameter) was filled with this solution and inserted into the cortex, and OGB1 and Alexa
183 were pressure-ejected from the pipette (–2 psi for 300–500 ms, 10–20 times). After
184 confirming loading, the craniotomy was sealed with a cover glass. 281.6 μm ×281.6 μm area
185 was imaged using a two-photon microscope (Nikon A1MP, Nikon), which was equipped
186 with a mode-locked Ti:sapphire laser (MaiTai Deep See, Spectra Physics) at 2 Hz with
187 512×512 pixels (0.55 μm /pixel). The excitation light was focused with a 25× Nikon (NA: 1.10)
188 PlanApo objective. The average power delivered to the brain was < 20 mW, depending on
189 the depth of focus. OGB1 and CTB555 were excited at 1000 nm except for the experiment
190 using 920 nm described in **Figure S1B**. The emission filters were 517–567 nm for OGB-1 and
191 600–650 nm for CTB555.

192

193 *Visual stimulation and image data acquisition*

194 The visual stimulation for studying the binocular response property was performed using
195 two LEDs controlled by Arduino (Smart Projects). To avoid contamination to the other eye,
196 LEDs were directly attached to the eyes with black silicon. Intensity was set as 40 cd/m². Each
197 stimulus started with a blank period (4s), which was followed by visual stimulation (4s). Each
198 stimulus was consisted of 4 rounds of alternative 0.5 seconds of ON and OFF periods. We
199 used 410 nm LED stimuli, based on the report that all mouse photoreceptors display similar
200 sensitivity to spectrum around 400 nm (Govardovskii et al., 2000). Our measures of
201 sensitivity are thus not biased towards responses originating from any specific
202 photoreceptor. In particular, the dorsal-ventral gradient in cone-opsin expression in the
203 mouse retina (Applebury et al., 2000; Sterratt et al., 2013) is not likely to have influence with
204 our measures of relative sensitivity. To study orientation and direction selectivity, drifting
205 square-wave gratings (100% contrast, 2 Hz temporal frequency) were presented in 12
206 directions of motion with 30 degree steps on a 27-inch LCD monitor (Samsung, Hwaseong,
207 South Korea) placed in 20 cm distance from eyes. We positioned the center of the monitor
208 to cover the vertical meridian. The drifting gratings visual stimulus sets were generated
209 using custom-made software written in PsychoPy (Peirce, 2007). The spatial frequency was
210 set at 0.03 cycles per degree. Each stimulus started with a blank period of uniform gray with
211 the same mean luminance of gratings (4s), which was followed by visual stimulation (4s).
212 For mapping of spatial frequency (SPF) and temporal frequency (TF) tunings, drifting sine-
213 wave gratings (100% contrast) were used. For SPF mapping experiments, sine-wave
214 gratings with six SPFs between 0.01 cycle per degree (cpd) and 0.32 cpd in octave steps,

215 drifting at 2 Hz were used. For TF mapping experiments, sine-wave gratings having 0.04 cpd
216 and drifting at 5 different TFs between 0.5 and 8 Hz in octave steps were used. Each stimulus
217 started with a blank period of uniform gray (4 s) followed by the same period of visual
218 stimulation during vertical and horizontal gratings were presented for 1 s for each of four
219 directions (0°, 180°, 90°, and 270° in that order). These 3 drifting grating stimulus sets were
220 repeated 10 times.

221 *Slice experiments*

222 Coronal slices (300 μ m thick) containing visual cortex were prepared from mice at postnatal
223 day 32–36 following deep anesthesia with isoflurane. At this age, binocular matching of
224 orientation preference in V1 neurons reaches the adult level, suggesting that binocular
225 visual functions in V1 are mature (Wang et al., 2010). Prior to recording, slices were
226 incubated for 1 h in oxygenated (95% O₂ and 5% CO₂) normal artificial cerebrospinal fluid
227 (ACSF) containing (in mM) 126 NaCl, 3 KCl, 1.3 MgSO₄, 2.4 CaCl₂, 1.2 NaH₂PO₄, 26 NaHCO₃,
228 and 10 glucose at 33°C as described previously (Ishikawa et al., 2014). Neurons with
229 pyramidal shaped somata in L2/3 were targeted by patch pipettes under fluorescent and
230 infrared differential interference contrast (DIC) optics with a 60X water immersion objective
231 (BX51, Olympus, Tokyo, Japan, **Figure 3A**). Pyramidal neurons retrogradely labeled with
232 CTB488 or CTB555 were identified by epifluorescence with appropriate filter sets (U-MWIB3:
233 excitation 479–495 nm and emission 510IF nm for CTB488; U-MWIG3: excitation 530–550
234 nm and emission 575IF nm for CTB555). We performed whole-cell recordings from pairs of
235 L2/3 pyramidal neurons at room temperature. In all three groups (CPN-CPN pairs, CPN-non
236 CPN pairs, and non-CPN pairs), we recorded from pairs of neurons with somata separated
237 within approximately 70 μ m. We analyzed only neurons showing regular spiking firing

238 patterns with a transient high-frequency firing at an initial phase followed by a low-
239 frequency firing for the rest of the response (**Figure S5A**). Cell bodies of the recorded
240 neurons were located $>50\ \mu\text{m}$ below the cut surface of the slice. The patch pipettes (4–6
241 $\text{M}\Omega$) were filled with an internal solution containing (in mM) 130 K-gluconate, 8 KCl, 1 MgCl_2 ,
242 0.6 EGTA, 10 HEPES, 3 MgATP, 0.5 Na_2GTP and 10 Na-phosphocreatine (pH 7.3, adjusted
243 using KOH). The presence of synaptic connections between recorded neuron pairs was
244 tested by applying brief depolarizing voltage pulses (duration, 1–2 ms) to evoke an action
245 potential in one cell (≥ 50 trials) and recording excitatory postsynaptic currents (EPSCs) from
246 the other cell (Ishikawa et al., 2014). Connection probabilities were calculated as the number
247 of unidirectionally or reciprocally connected neuron pairs over the number of recorded
248 neuron pairs. In the voltage-clamp mode, a brief, large depolarization pulse applied to the
249 soma initiates an action potential at the axon because of the imperfect space clamp. The
250 membrane potential of recorded cells was held at the reversal potential of IPSCs ($-70\ \text{mV}$)
251 after correction of liquid junction potential. Recordings were made using a Multiclamp 700B
252 amplifier, Digidata 1440A converter, and pCamp10 software (all from Molecular Devices).

253 Data were digitized at 10 kHz, filtered at 1 kHz, and analyzed using custom-made
254 programs written in MATLAB. To obtain the amplitude of EPSCs, we averaged the peak
255 values of individual EPSCs after excluding failures. The baseline current was defined as the
256 averaged current in a 5-ms window before application of depolarizing voltage pulses to the
257 presynaptic cell. The peak EPSC was defined as the peak current minus the baseline. We
258 selected cells with seal resistance $>1\ \text{G}\Omega$ and series resistance $<35\ \text{M}\Omega$ for analysis. For
259 measuring intrinsic membrane properties of neurons, 1-s step currents from $-100\ \text{pA}$ to 500
260 pA in 25 pA increments were injected in current clamp mode. The current injection

261 experiments were conducted within 5 min after establishing the whole-cell patch-clamp
262 configuration.

263

264 **QUANTIFICATION AND STATISTICAL ANALYSIS**

265 *Data analysis of visual response*

266 The images were analyzed by using our custom-written in-house software running on
267 MATLAB (Mathworks) as previously described (Hagihara et al., 2015; Ohki et al., 2005). In
268 brief, the cell outlines were automatically identified using template matching. The
269 identified cell outlines were visually inspected, and the rare but clear errors were manually
270 corrected. The time courses of individual cells were extracted by summing the pixel values
271 within the cell outlines. Slow drift of the time courses over minutes, potentially caused by
272 gradual z-movement, bleaching, and/or diffusion of OGB1, was removed by applying a
273 low-cut filter (cut-off, 2–4 min). The baseline (F) for each trial with each cell was calculated
274 by averaging the values of the last 0.5 s of the blank periods for all stimuli. Neuropil
275 contamination was removed as previously described (Hagihara et al., 2015; Kerlin et al.,
276 2010). Visually responsive cells were defined by ANOVA ($P < 0.01$) across blank and 2 eyes
277 (for ocular dominance stimuli), across blank and 12 direction periods (for
278 orientation/direction stimuli), across blank and 6 SPF/TF periods (for SPF/TF stimuli), where
279 $\Delta F/F > 2\%$. Instead of $P < 0.05$, $P < 0.01$ was used here, to reduce false positives due to large
280 sample size of neurons.

281 For responsive cells with ocular dominance stimuli, ODscore (Mrsic-Flogel et al.,
282 2007) was defined as follows: (response to ipsilateral-eye)/(response to ipsilateral-eye) +

283 (response to contralateral-eye). For responsive cells with drifting grating stimuli, as an index
284 for orientation selectivity tuning, gOSI was calculated. This value is equivalent to $:(\Sigma R(\theta_i)\sin(2\theta_i))^2 + (\Sigma R(\theta_i)\cos(2\theta_i))^2)^{1/2}/\Sigma R(\theta_i)$, where θ_i is the orientation of
285 each stimulus and $R(\theta_i)$ is the response to that stimulus. For direction selectivity, DI was
286 calculated as $:1 - (\text{response to null direction})/(\text{response to preferred direction})$. The
287 preferred orientation and direction were defined by vector averaging of responses ($\Delta F/F$)
288 to all orientations and directions, respectively. For correlation analysis, Pearson's correlation
289 coefficient was calculated to obtain pair-wises response correlation. Signal correlation was
290 calculated as the correlation coefficient between the trial-averaged traces in responses to
291 the ocular dominance stimulus. Noise correlation was obtained by subtracting the trial-
292 averaged traces from the traces to each trial, and then calculating the correlation coefficient
293 between those mean-subtracted traces. Note that the same analysis based on responses to
294 drifting gratings data gave consistent results (**Figure S4C, D**).

296

297 *Statistical analyses*

298 All data are expressed as the mean \pm standard error of the mean (SEM), unless stated
299 otherwise. The two-sided Wilcoxon signed-rank test was used to compare two paired
300 groups. Friedman test was performed when more than two groups were compared. When
301 significant, it was followed by Tukey-Kramer method. For distribution of preferred
302 orientation and direction, Kolmogorov-Smirnov test was used. For data from slice whole-
303 cell patch clamp recordings, Wilcoxon rank-sum test was used to compare two groups.
304 Fisher's exact test was used to compare the proportions of connected cell pairs. Kruskal-
305 Wallis test was used when more than two groups were compared. The variances between

306 groups were assumed to be similar. Throughout the study, $P < 0.05$ was considered
307 statistically significant, other than the definitions of visually responsive and selective cells
308 (see Data analysis of visual response). No statistical methods were used to pre-determine
309 sample sizes, but our sample sizes are similar to those generally employed in the field.

310

311 **DATA AND SOFTWARE AVAILABILITY**

312 All data and analysis code are available upon reasonable request to the Lead Contact, Kenta
313 M. Hagihara (hagi_k@med.kyushu-u.ac.jp; kenta-hagihara@umin.ac.jp).

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Results

To compare the visual response properties of CPNs and non-CPNs in L2/3 of mouse visual cortex, we performed *in vivo* two-photon calcium imaging (Ohki et al., 2005). Using visual stimulation to each eye, we first assessed the ocular dominance response properties of CPNs and non-CPNs. The fluorescent retrograde tracer cholera toxin B subunit conjugated to the fluorophore Alexa 555 (CTB555) was used to label CPNs (**Figure 1A, B, Methods**). As Oregon Green BAPTA-1 (OGB1) injection was also performed to largely cover the CTB555-positive region along the medial–lateral axis, our recordings equally contained binocular regions of V1 and lateral secondary visual cortex (more specifically LM (Wang and Burkhalter, 2007)). We imaged 1,866 CPNs and 11,399 non-CPNs from 7 mice. There was no difference in the fraction of visually responsive CPNs and non-CPNs ($31.3\% \pm 1.8\%$ vs. $28.9\% \pm 2.2\%$, $P = 0.30$, Wilcoxon signed-rank test, $n = 7$ mice). Note lower responsiveness than previous reports using drifting gratings (also see **Figure 1F**). Based on fluorescence signal changes in response to unilateral visual inputs, we calculated the ocular dominance (OD) score (see **Methods** and (Mrsic-Flogel et al., 2007)) for all visually responsive neurons. Surprisingly, some CPNs showed exclusive ipsilateral eye preference (**Figure 1C**). While OD scores varied extensively for both CPNs and non-CPNs, the proportion of neurons exhibiting a high OD score (with ipsilateral eye preference) was larger in CPNs than non-CPNs (**Figure 1D**). Indeed,

352 a significantly larger fraction of CPNs had OD scores >0.8 compared to non-CPNs (**Figure**
353 **1E**, $P = 0.016$, Wilcoxon signed-rank test, $n = 7$ mice), while there was no significant
354 difference in the fraction of OD scores <0.2 ($P = 0.47$). As previously reported in mice
355 (Antonini et al., 1999; Drager, 1974; Gordon and Stryker, 1996; Mrsic-Flogel et al., 2007; Scholl
356 et al., 2015), we did not observe any evidence for spatial clustering or microstructure related
357 to eye preference or to CPNs in C57BL/6 mice (**Figure S3**). Both CPNs and non-CPNs showed
358 similar tuning properties for drifting gratings, with only slightly greater orientation
359 selectivity in CPNs (**Figure 1F**, **Figure S4A, B**). These results suggest that CPNs have
360 somewhat distinct visual response properties from non-CPNs. Although non-CPNs could
361 include interneurons as well as pyramidal neurons with our current approach, we obtained
362 consistent results after correcting the potential influence by the small fraction of
363 interneurons (**Figure S4B**).

364 CPNs are not organized into columns or spatial clusters in mice (**Figure S3**), but it
365 is still possible that they form functional subnetworks. To test this possibility, we analyzed
366 the signal and noise correlations of CPN and non-CPN activity with unilateral eye
367 stimulation (**Figure 2**) and with drifting grating stimuli (**Figure S4D, E**). Both signal and
368 noise correlations were significantly stronger between CPN pairs than between CPN/non-
369 CPN pairs and non-CPN/non-CPN pairs (**Figure 2B**, CPN/CPN vs. CPN/non-CPN, $P < 0.001$;
370 CPN/CPN vs. non-CPN/non-CPN, $P < 0.001$; **Figure 2E**, CPN/CPN vs. CPN/non-CPN, $P < 0.001$;
371 CPN/CPN vs. non-CPN/non-CPN, $P < 0.001$; $n = 62$ planes from 7 mice; Friedman test
372 followed by Tukey-Kramer method). These differences also held true when we analyzed
373 neuron pairs of similar soma–soma distance (**Figure 2C and F**). Thus, CPNs are likely to form
374 local functional subnetworks that are organized independently of cell body location.

375 Pairs of V1 neurons showing stronger signal/noise correlation show higher
376 probability of synaptic connections (Cossell et al., 2015; Ko et al., 2011). To directly test
377 whether CPN pairs show higher connection probability, we conducted dual whole-cell
378 patch-clamp recordings from CPNs and/or non-CPNs in slices of the hemisphere
379 contralateral to the CTB injection site at 4–8 days post-injection, specifically targeting the
380 most densely CTB-labeled area in L2/3 (**Figure 3A, B**). We excluded putative interneurons
381 from the analysis of non-CPNs (**see Methods**). Single action potentials in a pyramidal
382 neuron produced EPSCs in a partner neuron in connected pairs (**Figure 3B**). The
383 latency of individual EPSCs was short (mean \pm s.d., 1.87 ± 0.49 ms) and the
384 fluctuation of the latency was low (0.2 ± 0.1 , coefficient of variation of the latency).
385 No reliable EPSCs with a latency longer than 2.52 ms was observed, indicating that
386 all the recorded EPSCs were monosynaptic currents. The probability of excitatory
387 connections between CPN/CPN pairs (30% of the recorded pairs) was significantly higher
388 than that between CPN/non-CPN pairs (12.2%, $P = 0.046$, Fisher's exact test, **Figure 3C**).
389 The probability of connections between non-CPN/non-CPN pairs (19.1%) also tended to
390 be lower than that of CPN/CPN pairs, although the difference did not reach significance.
391 The connection probability for all recorded pairs (21.1%, 28/133) was in agreement with
392 that for randomly sampled pairs in L2/3 visual cortex (Cossell et al., 2015; Ko et al., 2011).
393 We observed only one reciprocally connected CPN/CPN pair among all the recorded
394 pairs. The overall reciprocal connection probability (0.75%, 1/133) was similar to the
395 expected probability (1.19%, $(29/266)^2$), implying that the establishment of reciprocal
396 connections might be at the chance level regardless of projection targets of neurons
397 in the BZ. The amplitudes of EPSCs recorded from connected CPN/CPN pairs were

398 comparable to those measured from connected CPN/non-CPN and connected non-
399 CPN/non-CPN pairs ($P = 0.95$, Kruskal–Wallis test, **Figure 3D**). Also, there were no statistical
400 differences in intrinsic membrane properties between CPNs and non-CPNs (**Figure S5**).
401 Overall, these results demonstrate that L2/3 neurons sharing long-range projection targets
402 form functional subnetworks defined by local connectivity.

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420 **Discussion**

421 In the current study, we used CPNs as a model to investigate the functional properties and
422 local circuit organization of long-range projection neurons in the cortex. How CPNs and
423 other projection neurons in the early visual cortex distribute visual feature information in a
424 target-specific manner is an intriguing question. Our data showing stronger signal/noise
425 correlations (**Figure 2**) and local connectivity among CPNs (**Figure 3**) suggest the existence
426 of target-specific local functional subnetworks in L2/3. It has been reported that local
427 subnetworks of neurons sharing similar visual properties emerge through activity-
428 dependent synaptic plasticity (Ishikawa et al., 2014, 2018; Ko et al., 2013). Our data extend
429 these findings by showing that long-range projection specificity is another factor
430 determining local subnetwork organization in L2/3, a notion also supported by a recent
431 study that assessed V1 neurons projecting to different higher visual cortices (Kim et al.,
432 2018). In contrast to the strongly segregated subnetworks found in neurons projecting to
433 different higher-order visual areas within the same hemisphere (Kim et al., 2018),
434 moderately segregated subnetworks between CPNs and non-CPNs reported in the current
435 study may allow for interactions between the subnetworks of CPNs and other neurons in
436 the local network. Overall, this projection target-dependent segregation of local
437 subnetworks may be a general principle for intermingled projection neurons in the mouse
438 cortex. That said, different levels of segregation in distinct projection populations are of
439 note. The extent of segregation would define the selectivity of information to be extracted
440 locally and then transmitted to the projection target.

441 We found that a larger fraction of CPNs than non-CPNs preferentially responded
442 to ipsilateral eye input. Some of them even showed exclusive ipsilateral eye preference
443 (**Figure 1**). Considering that most of the retino-thalamic afferents cross in the chiasm

444 (Lund, 1965), and that most of the lateral geniculate neurons show binocular response in
445 rodents (Howarth et al., 2014), this strong ipsilateral selectivity found in CPNs was rather
446 unexpected. This ipsilaterally-biased CPN population may serve to duplicate visual
447 information conveyed by the thalamo-cortical projection, and send a copy of that
448 information to the other hemisphere (Laing et al., 2015). Thus, callosal connections may
449 function as a back-up system for the thalamo-cortical projection (Li et al., 2016). Different
450 from other projection neurons in V1 (Glickfeld et al., 2013), CPNs show similar SPF/TF
451 tuning as non-CPNs (**Figure S5**). This property may also be useful for the putative back-up
452 function. In contrast to our findings, a recent study (Lee et al., 2019) did not find a strongly
453 ipsilaterally biased population (to the hemisphere where CPN somata are located)
454 among callosal inputs to L2/3 V1 neurons (contralateral to the callosal recipient neurons),
455 raising the possibility that the ipsilaterally biased callosal population selectively targets
456 contralateral L5 neurons, which receive the largest fraction of callosal inputs (Petreanu et
457 al., 2007), whereas majority of callosal neurons, which show contralateral eye (to the
458 hemisphere where CPN somata are located) preference, might target contralateral L2/3
459 neurons.

460 In the current study, we did not address whether ipsilaterally-tuned CPNs (or
461 more in general, CPNs with similar response properties) selectively tend to show higher
462 connectivity than random CPNs. This is likely the case given the previous findings that
463 similarly tuned neurons tend to be connected in the monocular region of V1 (Cossell et al.,
464 2015; Ko et al., 2011) and that establishment of such fine-scale subnetworks relies on
465 correlated neuronal activity during development (Ishikawa et al., 2014; Ko et al.,
466 2013). It would be an intriguing future study to directly address this point by the post-

467 hoc slice recording method following *in vivo* imaging (Cossell et al., 2015; Ko et al.,
468 2013; Ko et al., 2011).

469 Several recent studies have shown that V1 neurons projecting to different
470 higher-order visual cortices (Glickfeld et al., 2013; Matsui and Ohki, 2013) and subcortical
471 areas show distinct response properties (Kim et al., 2015; Lur et al., 2016). However, the
472 extent of overlap through collateral projections (Han et al., 2018) is not known. For
473 instance, CPNs could overlap with other specific projecting populations (Yamashita et al.,
474 2013; Yamashita et al., 2018). Thus, comprehensive characterization of the relationships
475 among projection target, response properties, molecular identity, and local connectivity is
476 required for a better understanding of the circuit logic in the early visual cortex, which
477 functions as a central hub to process visual information.

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505 **Author Contributions**

506 K.M.H. and Y.T. initially conceived the research. K.M.H. designed and performed imaging
507 experiments and analyzed the data. K.O. supervised imaging experiments, data analysis,
508 and interpretation of the data. A.W.I designed and performed slice experiments and
509 analyzed the data. Y.Y. supervised slice experiments, data analysis, and interpretation of
510 the data. K.M.H., Y.T., A.W.I. wrote the manuscript. K.O. and Y.Y commented on the
511 manuscript.

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513 The authors declare no competing interests.

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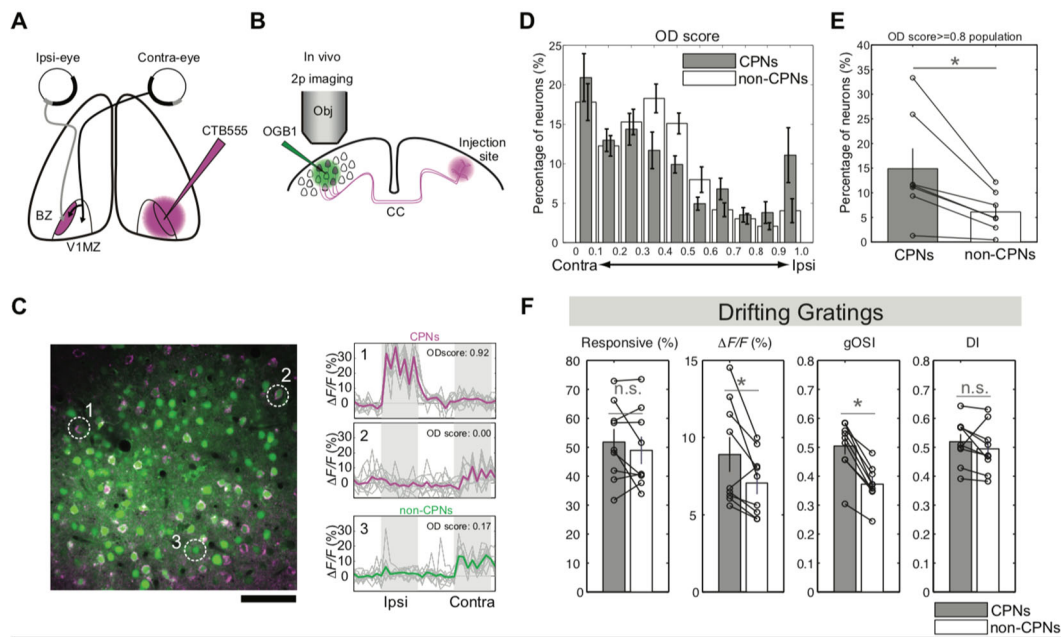
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688 **Figures**



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691 **Figure 1. Ocular dominance response properties of CPNs and non-CPNs**

692 **A.** Schematic drawing of CPNs labeling with retrograde tracer CTB555. BZ: binocular zone;
 693 V1MZ: primary visual cortex monocular zone.

694 **B.** Schematic drawing of *in vivo* two-photon calcium imaging of CTB555 labeled CPNs and
 695 surrounding non-CPNs.

696 **C.** (Left) Image of OGB1-labeled (green) and retrogradely CTB555-labeled (magenta) callosal
 697 projection neurons (CPNs) and non-CPNs in the binocular zone (BZ). Scale bar: 20 μ m.
 698 (Right) Representative calcium traces from CPNs (#1 and #2) and a non-CPN (#3) shown in
 699 the left panel.

700 **D.** Distribution of ocular dominance (OD) scores for CPNs and non-CPNs.

701 **E.** Proportion of neurons with an OD score over 0.8. * $P < 0.05$; (n = 7 mice, Wilcoxon signed-
 702 rank test)

703 **F.** Responsiveness, $\Delta F/F$, gOSI, and direction index (DI) of CPNs and non-CPNs.

704 *: $P < 0.05$; n.s., non-significant; (Wilcoxon signed-rank test).

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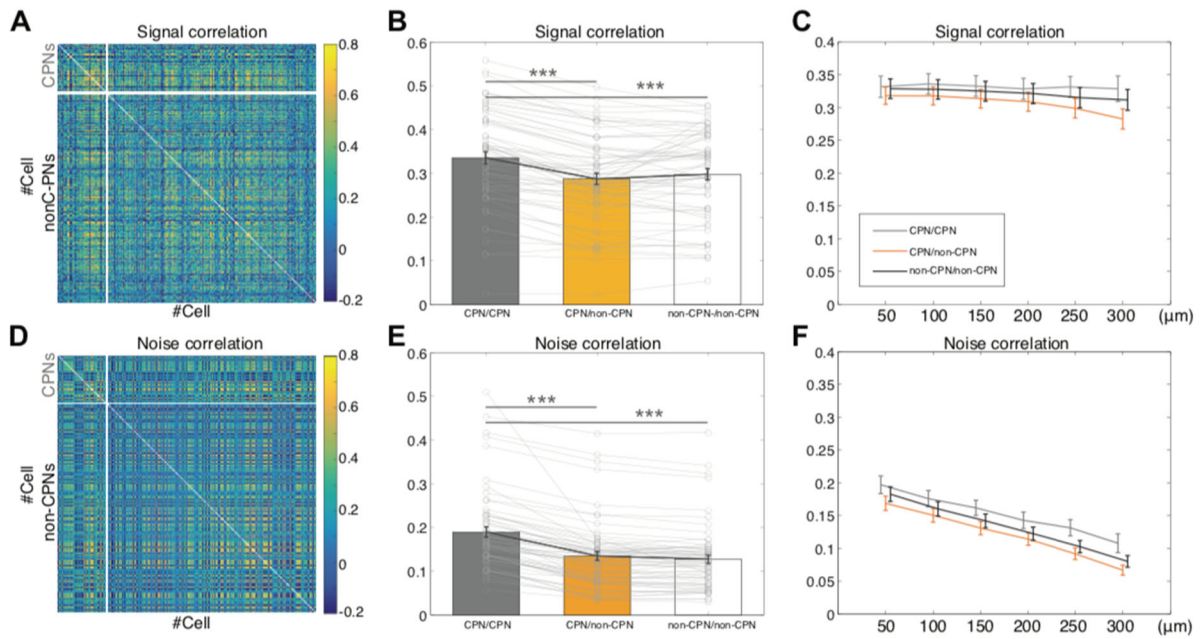
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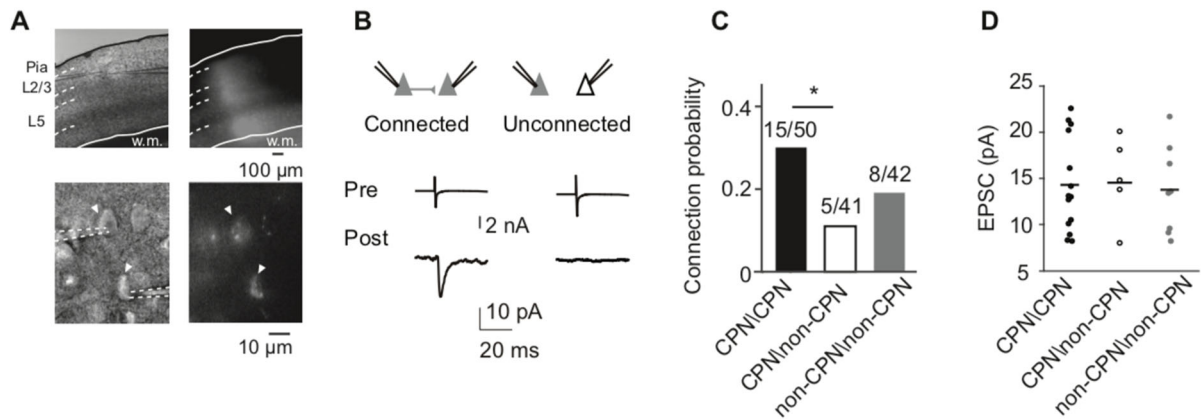
Figure 2. Correlation structure of CPNs and non-CPNs

A, D. Representative signal (**A**) and noise (**D**) correlation matrices of neurons from a single imaging plane. The order of neurons is sorted so that CPNs are located at the top and non-CPNs at the bottom.

B, E. Means of signal (**B**) and noise (**E**) correlation values from each plane. CPN/CPN pairs show both higher signal and noise correlation coefficient values than CPN/non-CPN and non-CPN/non-CPN pairs.

C, F. Signal (**C**) and noise (**F**) correlations of CPN/CPN (gray), CPN/non-CPN (orange), and non-CPN/non-CPN (black) pairs with respect to cortical distance.

***: $P < 1.0 \times 10^{-3}$; (Friedman test followed by Tukey-Kramer).



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737 **Figure 3. Local synaptic connectivity of CPNs**

738 **A.** Top-left: Differential interference contrast image showing a recording electrode position
 739 in L2/3. Top-right: Corresponding image of CTB fluorescence in a coronal slice of primary
 740 visual cortex containing the binocular zone. w.m.: white matter. Bottom-left: Higher
 741 magnification of the interference contrast image showing simultaneously recorded
 742 neurons (marked with white triangles). Recording pipettes were marked with dotted lines.
 743 Bottom-right: Corresponding high-magnification fluorescent image of CTB-labeled CPNs.
 744 White triangles mark the same CTB-labeled recorded neurons.

745 **B.** Example averaged traces of presynaptic action currents evoked by depolarizing voltage
 746 pulses and resultant EPSCs in postsynaptic cells.

747 **C.** Connection probabilities of CPN/CPN, CPN/non-CPN and non-CPN/non-CPN pairs. The
 748 connection probability of CPN/CPN pairs was significantly higher than that of CPN/non-CPN
 749 pairs ($*P < 0.05$ from Fisher's exact test). The numbers of recorded and connected cell pairs
 750 are shown above each bar.

751 **D.** Amplitudes of evoked EPSCs obtained from connected neuron pairs. Black lines indicate
 752 medians.

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