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Direction and orientation preferences in mouse superior colliculus and its retinal inputs exhibit a topography of cardinal biases atop locally mixed tuning

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eLife Assessment

This is a potentially **important** work on the organization of visual information in the rodent superior colliculus. It reports that the selectivity of neurons to line orientation and motion in the visual image is largely governed by the sensitivities of retinal neurons and their ordered projection to the superior colliculus. If confirmed, these conclusions could substantially revise prior thinking in this field. However, in the present state, the methods and analysis are **incomplete** and cannot justify all the claims.

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Abstract

In mouse superior colliculus (SC), neurons are robustly tuned to motion direction and orientation, but it remains controversial whether these preferences form global maps or are arranged in a salt-and-pepper fashion, and how this organization relates to that found in the retinal input to the SC. We addressed these questions by combining two-photon calcium imaging of retinal boutons and SC neurons in superficial SC with Neuropixels recordings across the full depth of SC. We then asked how direction and orientation preferences depend on visual-field location and how tuning similarity among units depend on lateral and vertical distance in the SC. Retinal boutons were strongly tuned and their preferences closely matched the retinal topographic organization of cardinal motion directions and orientations previously described for direction- and orientation-selective ganglion cells, showing that SC receives a highly ordered, retinocentric code for motion and edges. With increasing depth, SC neurons progressively deviated from this organization. On top of this cardinal topography, local clustering of tuning preferences in retinal boutons and SC neurons was weak and confined to very small spatial scales. We did not detect robust global maps of direction or orientation. Together, these results show that mouse SC inherits a topographic organization of cardinal directions and orientations from the retina, progressively diverts from this orderly organization with increasing distance from the retinal input, and exhibits broad coverage of directions and orientations at each visual location, likely supporting flexible readout for diverse visually guided behaviors.

Introduction

Visual feature maps in the cortex can range from smoothly organized columns to largely disordered “salt-and-pepper” arrangements. In carnivores and primates, orientation and direction preferences in primary visual cortex (V1) form orderly maps (Basole et al., 2003 [DOI](#); Hubel and Wiesel, 1968 [DOI](#), 1962 [DOI](#); Ibbotson and Jung, 2020 [DOI](#)), whereas in rodent V1, similarly tuned

neurons are embedded in a largely salt-and-pepper background with only weak local clustering (Kaschube, 2014 [↗](#); Ohki et al., 2005 [↗](#)). While the SC is the major retinorecipient structure in the mouse midbrain and plays a central role in visually-guided orienting, escape, and prey capture (Hoy and Farrow, 2025 [↗](#); Wheatcroft et al., 2022 [↗](#)), its functional organization remains debated (Ahmadlou and Heimel, 2015 [↗](#); Chen et al., 2021 [↗](#); De Malmazet et al., 2018 [↗](#); Feinberg and Meister, 2014 [↗](#); Inayat et al., 2015 [↗](#); Kasai and Isa, 2021 [↗](#); Li et al., 2020 [↗](#)). Neurons in superficial SC (sSC) are robustly tuned to direction and orientation (De Franceschi and Solomon, 2018 [↗](#); Inayat et al., 2015 [↗](#); Ito et al., 2017 [↗](#); Wang et al., 2010 [↗](#)), but it is unclear whether their preferences form systematic maps across the visual field or are arranged in a largely mixed fashion.

Previous studies have reported conflicting organizations of direction and orientation in mouse SC. Some found strong clustering and global structure (Feinberg and Meister, 2014 [↗](#); Li et al., 2020 [↗](#)), including concentric orientation maps centered on the nose (Ahmadlou and Heimel, 2015 [↗](#); De Malmazet et al., 2018 [↗](#)) and a predominant motion direction that reverses near the binocular border (De Malmazet et al., 2018 [↗](#)). Other work reported weak direction clustering and no evidence for stereotypical direction columns (Chen et al., 2021 [↗](#); Inayat et al., 2015 [↗](#); Kasai and Isa, 2021 [↗](#)). These discrepancies suggest that the presence, strength, and spatial scale of direction and orientation maps in SC may depend on specific experimental conditions.

In parallel, work in the retina has revealed a highly structured, retina-centered topographic organization of motion and orientation preferences. Direction-selective and orientation-selective ganglion cells (DSGCs and OSGCs) align their preferred directions and orientations with longitudinal lines defined by body and gravity axes and latitudinal lines defined by two additional axes (Laniado et al., 2025 [↗](#); Sabbah et al., 2017 [↗](#); Tiriach et al., 2022 [↗](#); Vita et al., 2024 [↗](#)). Because most retinal ganglion cell (RGC) types, including all cardinal direction-selective types, project retinotopically to sSC (Ellis et al., 2016 [↗](#); Kay et al., 2011 [↗](#)), this topographic organization provides a concrete prediction for direction and orientation preferences at each collicular location. It is unclear how this structured retinal input is expressed at the level of retinal boutons in SC, and how it is transformed in SC neurons across the surface and with depth.

Here, we combined two-photon calcium imaging of retinal boutons and SC neurons in posterior sSC with Neuropixels recordings across the full depth of anterior SC. Using receptive-field (RF) mapping, we positioned each unit into visual space and compared preferred directions and orientations to predictions of the retinal topographic organization. We quantified local clustering and large-scale topographic structure of tuning and examined how alignment to the retinal organization changed with depth. Our results show that SC inherits a topographic organization of cardinal direction and orientation preferences at the level of retinal boutons, while tuning preferences in local populations of retinal boutons and SC neurons are locally mixed and only weakly clustered.

Results

Retinal boutons and sSC neurons are robustly tuned to motion direction and orientation with shared cardinal biases

To compare the functional organization in retinal inputs and SC neurons, we imaged calcium signals in synaptic boutons of RGC axons and in neurons in the posterior right SC. In one group of mice, we expressed a synaptically targeted calcium indicator (Dreosti et al., 2009 [↗](#)) in RGCs by intravitreal injection of AAV2-SyGCaMP6f into the left eye (Figure 1A [↗](#)). In another group of mice, we expressed a cytosolic calcium indicator (Chen et al., 2013 [↗](#)) in neurons in the superficial layers of the right SC by injecting AAV1-GCaMP6f (Figure 1C [↗](#)). To expose the posterior SC representing the top-left peripheral visual field, we used a cylindrical cranial implant to displace the transverse sinus without damaging cortex. Mice were head-fixed on a treadmill during two-photon imaging, and recorded movies of calcium signals were motion-corrected and segmented into regions of interest (ROIs) with an adapted version of Suite2p (Pachitariu et al., 2016 [↗](#)) that accounted for lateral and axial brain motion.

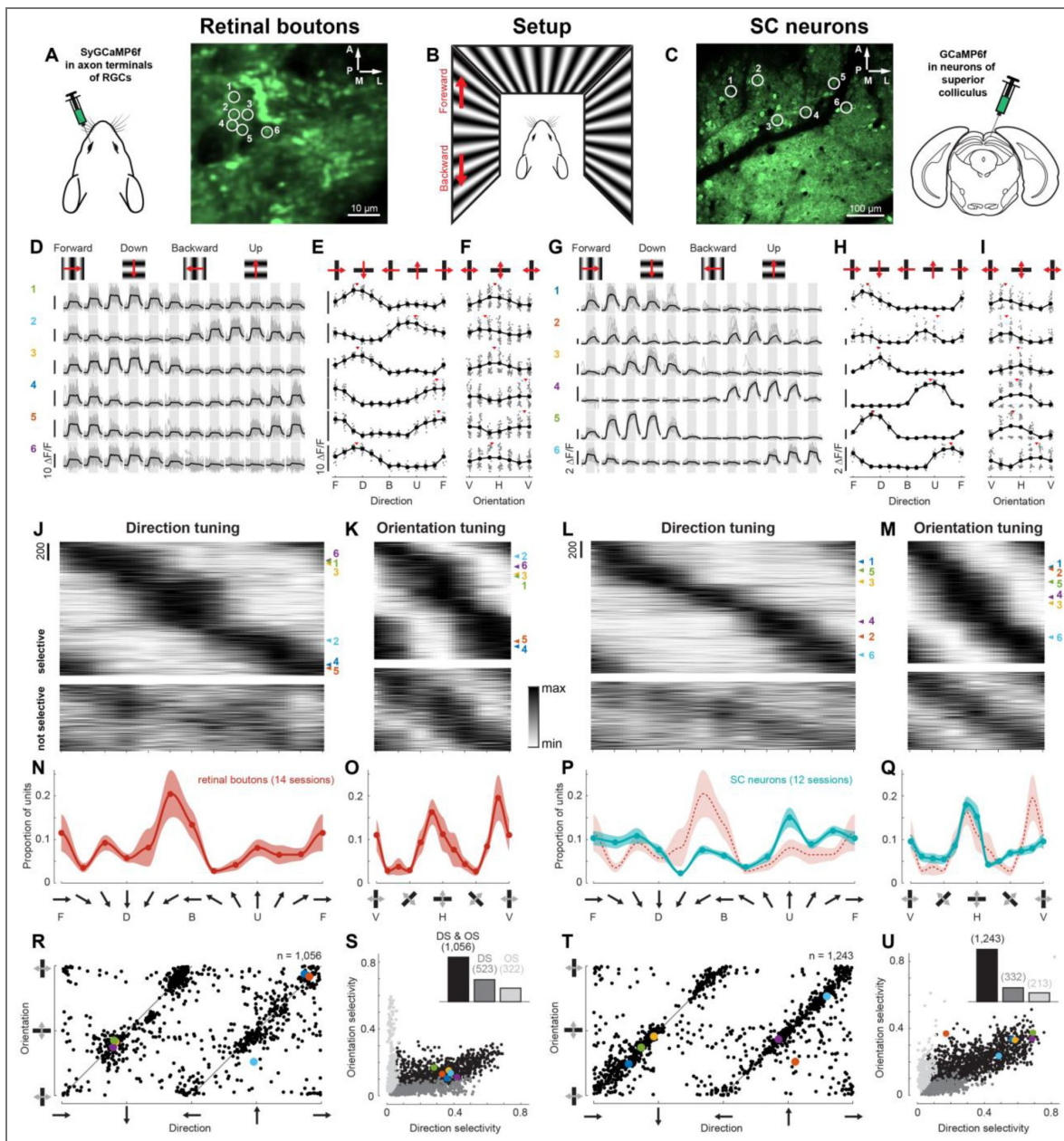


Figure 1. Retinal boutons and neurons in the SC were well tuned to motion direction and orientation.

A. AAV-SyGCaMP6f was delivered to RGCs via intravitreal injection (left). Example average image of retinal boutons in the SC (right). B. Visual stimuli were presented on three monitors covering the visual field from -135° to 135° azimuth and -42° to 42° elevation. Grating stimuli covered all screens. Stimulus directions—specifically “forward”/temporo-nasal and “backward”/naso-temporal—were described from the perspective of the left eye, because RFs of all recorded units were in the left hemifield. C. AAV-GCaMP6f is delivered to SC neurons via injection (right). Example average image of neurons in the SC (left). D. Single-trial fluorescence traces (gray) and mean fitted traces (black) of example boutons (marked in A) in response to sinusoidal gratings drifting in 12 directions. Gray shading indicates the 2 s stimulus presentations. E–F. Direction (E) and orientation (F) tuning curves of example boutons (same as in D), showing single-trial response amplitudes (gray dots), mean amplitudes (black dots), and preferred direction/orientation (red triangles). F – forward, D – down, B – backward, U – up, V – vertical, H – horizontal. G–I. As in D–F, for example SC neurons (marked in C). J–M. Direction (J) and orientation (K) tuning curves of all DS or OS (top) and non-selective (bottom) retinal boutons recorded across all sessions, sorted by their preferred direction of motion or orientation. Each tuning curve was scaled to a range between 0 and 1. Triangles indicate tuning curves of example boutons from A. Corresponding data for SC neurons in L and M. N–Q. Mean distribution ($\pm$ SEM) of preferred directions (N) and orientations (O) across all sessions of retinal boutons (red). Datasets with fewer than 20 selective units were excluded. Corresponding data for SC neurons (teal) in P and Q. For comparison, mean distributions of retinal boutons are overlaid (red; same data as in N, O). R. Preferred direction versus orientation for retinal boutons selective to both features. Example boutons from A marked in color. S. Direction versus orientation selectivity for all tuned retinal boutons. Example boutons from A marked in color. Insets show histograms of selectivity types. T, U. As in R and S, for SC neurons.

We measured tuning to direction of motion and orientation of drifting sinusoidal gratings. Visual stimuli were presented on three monitors surrounding the animal, covering -135° to 135° azimuth and -42° to 42° elevation (Figure 1B). All stimuli were first defined in spherical visual coordinates and then mapped onto the flat monitor surfaces. Gratings moved in 12 directions (30° steps) at a fixed spatial frequency of 0.08 cycles/ $^\circ$, temporal frequency of 2 cycles/s, and 100% contrast. For each bouton and neuron, we fitted responses with a single temporal kernel aligned to stimulus onset and an independent amplitude per trial, separating responses to the current stimulus from residual activity induced by the previous stimulus. This model captured responses well (Figure 1D,G; mean $\pm$ standard deviation (SD) explained variance 0.32 ± 0.20 for boutons, 0.44 ± 0.23 for neurons), and identified 65% of retinal boutons (2,391/3,677) and 50% of SC neurons (2,378/4,769) as responsive to drifting gratings. Based on the fitted amplitudes, we calculated the vector average over all motion directions and all orientations. For both direction and orientation, preference and selectivity were defined as the angle and length of the resulting vector (Figure 1E,F,H,I). This measure of orientation tuning has also been termed axis tuning, because it is based on responses to motion along an axis (opposite directions combined) rather than to static stimuli. Orientation preferences were consistent across moving and static stimuli (Figure S1K,L), so we will use the more common term *orientation selectivity*. A bouton or neuron was considered direction or orientation tuned if its selectivity was significantly larger than zero.

Retinal boutons and SC neurons were frequently selective to direction or orientation and exhibited a shared bias toward cardinal axes. Among responsive boutons, 23% were direction-selective (DS) only, 13% orientation-selective (OS) only, 44% selective to both features, and 20% non-selective (Figure 1S). Responsive SC neurons comprised a larger fraction of units selective to both direction and orientation (52%), fewer units selective to a single feature (DS-only: 14%; OS-only: 9%), and more non-selective units (25%) (Figure 1U). In a subset of SC neurons, we also presented moving bars and static gratings and found that preferred directions and orientations were highly consistent, confirming that tuning was largely invariant to stimulus class (Figure S1A-L) (Ahmadlou and Heimel, 2015; Inayat et al., 2015). Across both retinal boutons and SC neurons, preferred directions and orientations spanned the full range, but were overrepresented near cardinal angles (0° , 90° , 180° , 270° ; Figure 1J-M), which was consistent across recordings (Figure 1N-Q) and in line with previous reports in RGCs (Baden et al., 2016; Kerschensteiner and Hardesty, 2022; Laniado et al., 2025; Nath and Schwartz, 2016; Sabbah et al., 2017; Tiriach et al., 2022; Vita et al., 2024) and SC neurons (Chen et al., 2021; Inayat et al., 2015). Preferred directions strongly predicted preferred orientations (Figure 1R,T), and median direction selectivity exceeded orientation selectivity (boutons: 0.21 vs 0.13; neurons: 0.24 vs 0.17; Figure 1S,U, Figure S1M,N).

A direct comparison between retinal boutons and SC neurons revealed subtle differences in tuning. Both boutons and neurons showed cardinal biases, but boutons displayed a stronger bias toward naso-temporal motion (consistent with optic flow during forward locomotion), while SC neurons were more biased toward near-horizontal orientations (Figure 1N-Q). Median direction selectivity was similar across populations (0.21 for boutons vs 0.24 for neurons; $p = 0.11$, Wilcoxon rank-sum test; Figure S1M), whereas median orientation selectivity was significantly higher in neurons (0.13 vs 0.17; $p < 0.001$, Wilcoxon rank-sum test; Figure S1N). Together, these results show that retinal boutons and SC neurons in posterior SC share robust direction and orientation tuning with cardinal biases, but differ in the balance between directional bias and orientation selectivity.

Superficial SC inherits retinal topography of cardinal direction and orientation preferences

Elegant studies in mouse retina have shown that tuning preferences of DSGCs and OSGCs are systematically organized in retina-centered spherical coordinates (Laniado et al., 2025; Sabbah et al., 2017). DSGCs form four main cardinal classes whose preferred motion directions at each retinal location align with longitudinal lines defined by the body axis and the gravitational axis, corresponding to optic flow during forward, backward, upward and downward self-motion

(Sabbah et al., 2017 [↗](#)) (Figure 3A [↗](#)). OSGCs form an analogous spherical topography, in which preferred orientations align either with one of two longitudinal fields or with two latitudinal fields, that is, concentric lines around one of two axes (Laniado et al., 2025 [↗](#)) (Figure 3B [↗](#),C [↗](#)). These retinal studies thus provide a geometric model that predicts, for any given RF position in visual space, which motion directions and edge orientations are preferred by RGCs.

To relate our SC data to this topographic organization of the retina, we first mapped RFs of retinal boutons and SC neurons using sparse noise (Figure 2A–F [↗](#)). We asked how well RF locations followed retinotopy by fitting, for each imaging field of view (FOV), a simple linear retinotopic map that optimally translated, scaled, and rotated positions in the FOV to best match RF locations in visual space. Mapped RF positions deviated from the positions predicted by this fit by only 2.63° for boutons and 2.93° for neurons on average, corresponding to roughly one quarter of the RF diameter (Figure 2G–J [↗](#)). RFs could be mapped for the majority of direction- or orientation-tuned units (boutons: 85% and 86%; neurons: 66% and 67%). The proximity of RFs to the monitor edges did not systematically affect measured direction or orientation preferences (Liang et al., 2023 [↗](#); Roth et al., 2018 [↗](#)) (Figure S2 [↗](#)). Nevertheless, we excluded units with RF centers closer than 5° to any monitor edge from further analyses.

The retinal topographic organization of directions and orientations was largely preserved in retinal boutons and partially preserved in SC neurons. For each unit, we projected its RF location onto the axes of the retinal geometric model (Figure 3A–C [↗](#)) and compared its measured preferred direction and orientation to the corresponding predicted vectors (Figure 3D–G [↗](#)). In retinal boutons, the median angular difference between the measured preferred direction and the closest of the four predicted directions was 11.6°, smaller than expected for a model with a uniform distribution of preferences (21.0–24.0°; Figure 3D [↗](#),L [↗](#)). The median difference was also smaller than in a null distribution obtained by permuting RF locations (13.9–15.4°; Figure 3L [↗](#)), showing that direction preferences varied systematically with RF position. For orientation preferences in boutons, the median difference was 7.5°, again smaller than in the permuted (8.8–10.1°) and uniform (14.1–16.9°) controls (Figure 3E [↗](#),M [↗](#)). Direction preferences of SC neurons matched the model to a similar extent as boutons ($p = 0.73$, Wilcoxon rank-sum test on angular differences), with a median difference of 12.2°, smaller than in permuted (13.3–14.5°) and uniform (21.0–24.0°) controls (Figure 3F [↗](#),N [↗](#)). In contrast, orientation preferences of SC neurons deviated more strongly from the geometric predictions than those of boutons ($p < 1e-4$, Wilcoxon rank-sum test): the median difference of 10.1° was comparable to the permuted control (9.9–11.2°), although still smaller than the uniform model (13.2–15.7°; Figure 3G [↗](#),O [↗](#)). Restricting the analysis to DS-only or OS-only units yielded the same qualitative picture (Figure S3 [↗](#)). These findings show that retinal boutons in posterior SC closely preserve the retinal topography of direction and orientation tuning, while SC neurons preserved the topography for direction but deviated from that for orientation.

Retinal boutons were biased toward temporal motion-preferring cells, whereas SC neurons showed a more uniform distribution across direction classes. We assigned each tuned bouton or neuron to the predicted direction or orientation vector closest to its measured preference. For direction, we used the four optic-flow classes defined by Sabbah et al. (2017) [↗](#), for example “advance cells”, which prefer optic flow in naso-temporal direction elicited by forward self-motion (Figure 3A [↗](#)). Retinal boutons were strongly biased toward the “advance” class (40% of direction-tuned boutons, Figure 3H [↗](#)), consistent with their stronger bias for backward/naso-temporal stimulus motion in the overall direction distribution (Figure 1N [↗](#),V [↗](#)) and with the bias found in DSGCs (Sabbah et al., 2017 [↗](#)). SC neurons, by contrast, were more evenly distributed across direction classes, with 19–28% of direction-tuned neurons in each class (Figure 3J [↗](#)). For orientation, boutons were roughly balanced between vertical and horizontal classes (52% vs 48%; Figure 3I [↗](#)), while SC neurons were slightly biased toward horizontal orientation classes (65%; Figure 3K [↗](#)). Restricting the analysis to DS-only or OS-only units, however, yielded slightly different distributions (Figure S3E–H [↗](#)).

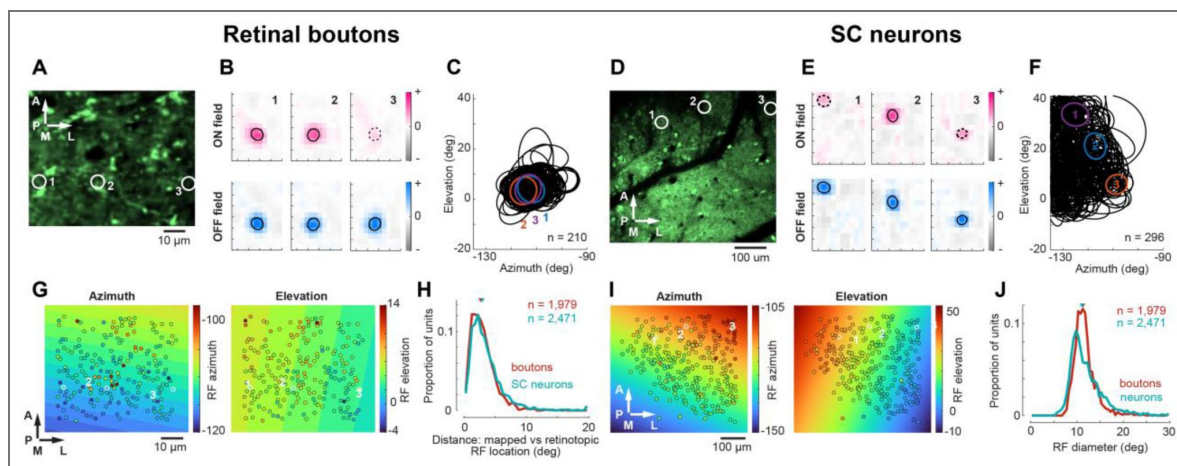


Figure 2. RF locations deviated from retinotopy by about one quarter of their diameter.

A. Example average image of retinal boutons in the SC. B. ON (top) and OFF (bottom) receptive subfields of 3 example retinal boutons (marked in A). Outlines (center $\pm$ 1 SD of fitted 2D Gaussian) are shown in black. Examples 1 and 2 had significant ON and OFF subfields, while example 3 had only an OFF subfield (dashed outline of the ON subfield is shown for reference). C. RF outlines for all retinal boutons recorded in session from A. Example boutons marked in color. D–F. Same as in A–C, for an example recording of SC neurons (same FOV as in Figure 1C). G. Relationship between position in the FOV and RF azimuth (left) or elevation (right) for retinal boutons from the session in A. The colored background gradients represent the best linear fit between FOV and RF positions. H. Distances between mapped and retinotopic (predicted from FOV position) RF locations for retinal boutons (red) and SC neurons (teal). Retinal boutons had slightly smaller distances than SC neurons (medians (triangles): 2.6° and 2.9° for boutons and SC neurons; $p < 1e-6$, Wilcoxon rank-sum test). I. Same as in G, for SC neurons and example session from D. J. RF diameters (2 SDs of the fitted Gaussian) across retinal boutons (red) and SC neurons (teal). RF sizes were not significantly different between boutons and SC neurons (medians (triangles): 11.3° and 11.1° for boutons and SC neurons; $p = 0.11$, Wilcoxon rank-sum test).

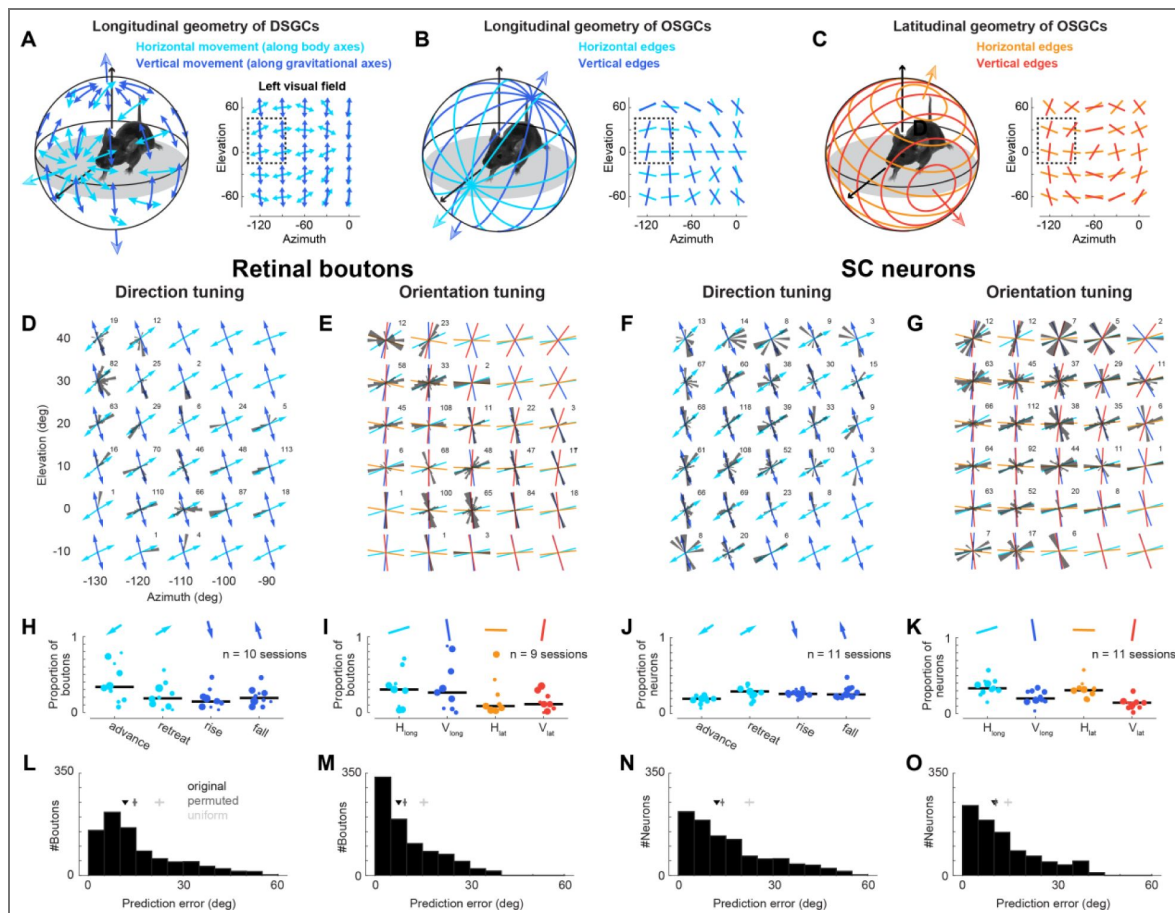


Figure 3. Topography of directions and orientations was consistent with topography originally identified in the retina.

A–C. Cartoons illustrating the topography of direction preferences (A) of DSGCs and orientation preferences (B, C) of OSGCs for the left eye. Direction preferences align with two axes defining longitudes (A), while orientation preferences align with two axes defining longitudes similar to those for direction (B) and two further axes defining latitudes (C). Dotted rectangle within left visual field marks the part of the visual field that contains RFs of our data (shown in D–G). D–G. Direction (D) and orientation (E) preferences of retinal boutons across visual space (RF positions based on mapped RFs, not retinotopy). Arrows in D point into directions defined by the DSGCs longitudes (A). Colored lines in E follow orientations of longitudes (B) and latitudes (C) for OSGCs. Numbers indicate unit count in each polar histogram. Corresponding data for SC neurons in F and G. RF positions for SC neurons were based on retinotopic fits (see Figure 2). H–K. Classification of all DS (H) and OS (I) boutons according to the closest predicted vector from the geometric model. Each dot shows one recording session (dot size reflects total number of boutons); black line indicates median proportion across sessions. Naming follows previous conventions, where DSGC classes were named after the behavior that elicits their responses, e.g., “advance cells” for DSGCs preferring movement in naso-temporal direction. Arrows and lines at the top are example vectors from a central location in D and E. Corresponding data for SC neurons in J and K. L–O. Angular differences between preferences and closest predicted vectors for direction (L) and orientation (M) across all tuned retinal boutons. The median difference (triangle) of the original data is compared to the median (2.5–97.5 percentile intervals) expected when RF locations were randomly permuted (dark gray) or when preferences were randomly drawn from a uniform distribution (light gray). Direction: 11.6° original, 13.9°–15.4° permuted, 21.0°–24.0° uniform; orientation: 7.5° original, 8.8°–10.1° permuted, 14.1°–16.9° uniform. Corresponding data for SC neurons in N and O. Direction: 12.2° original, 13.3°–14.5° permuted, 21.0°–24.0° uniform; orientation: 10.1° original, 9.9°–11.2° permuted, 13.2°–15.7° uniform.

Superficial SC and its retinal input lack global maps of direction or orientation

Smoothed maps of population tuning across visual space lacked systematic gradients, and only retinal boutons showed local tuning consistency above chance. While the previous analyses have revealed a topography of cardinal direction and orientation preferences, we next asked whether there are local biases towards single directions or orientation that could form global maps in the sSC. Visualizing preferred directions and orientations at RF locations across all recordings revealed mixtures of preferences at essentially all positions (Figure 4A–D). Averaging preferences within circular patches of 10° diameter to construct smoothed maps across visual space likewise revealed no smooth gradients or concentric patterns in either boutons or neurons (Figure 4E–H). We then quantified local consistency as the inverse circular variance of preferences within each patch (Figure 4I–L). Retinal boutons showed substantially higher local consistency than expected from null distributions obtained by permuting RF locations (direction: 0.48 ± 0.001 ; orientation: 0.52 ± 0.001 ; mean $\pm$ standard error of the mean (SEM); Figure 4I,J,M,N), and in most individual bouton recordings both direction and orientation consistency exceeded the shuffled control (Figure 4Q,R). In SC neurons, by contrast, local consistency was lower (direction: 0.23 ± 0.0003 ; orientation: 0.33 ± 0.0003 ; mean $\pm$ SEM; Figure 4K,L) and indistinguishable from null distributions for both features (Figure 4O,P). No recording showed elevated direction consistency and only three recordings showed elevated orientation consistency (Figure 4S,T). For both retinal boutons and SC neurons, consistency was not trivially explained by the number of units per patch (Figure S4). The somewhat higher tuning consistency in local populations of retinal boutons may be explained by subsets of retinal boutons originating from the same RGC and therefore sharing identical tuning. More broadly, the absence of a consistent direction and orientation map in sSC and the relatively weak tuning consistency among boutons and neurons contrasts previous findings, which would predict a clear overrepresentation of near-vertical orientation preferences and a mix of backward and upward direction preferences in the posterior SC (Ahmadlou and Heimel, 2015; De Malmazet et al., 2024, 2018; Feinberg and Meister, 2014; Li et al., 2020); these discrepancies are addressed in the Discussion.

Local functional clustering in sSC is weak and limited to very small spatial scales

Pairs of retinal boutons showed more similar tuning preferences than pairs of SC neurons. In example FOVs, preferences of boutons and neurons showed intermingled directions and orientations without obvious clusters (Figure 5A–D). When we considered all pairs of simultaneously recorded units, pairwise differences were smaller than expected for a uniform distribution of preferences (90° for direction, 45° for orientation) but still large: mean differences were ~75° for direction and 35° for orientation among boutons, and ~85° and 44° among neurons (Figure 5E–H).

Pairwise tuning differences showed only weak dependence on lateral distance and departed from chance levels only for very close pairs. Pairwise differences as a function of lateral distance in the SC closely followed null distributions obtained by randomly permuting unit locations within each FOV (Figure 5E–H). For retinal boutons, only pairs separated by $\leq 10 \mu\text{m}$ had direction and orientation preferences that were clearly more similar than expected from the null distribution (Figure 5E,F). For SC neurons, elevated similarity was restricted to pairs within $< 20 \mu\text{m}$, with orientation differences remaining slightly below the null distribution up to $\sim 180 \mu\text{m}$ but with a very small absolute deviation ($< 4^\circ$; Figure 5G,H). Across recordings, this weak clustering at very short distances was robust for boutons but rare for neurons: most bouton recordings contained nearby pairs with tuning differences below the null distribution, whereas only few neuron recordings did so (Figure 5I–L). Because bouton recordings covered smaller FOVs than neuron recordings, we tested whether FOV size could explain the higher similarity among

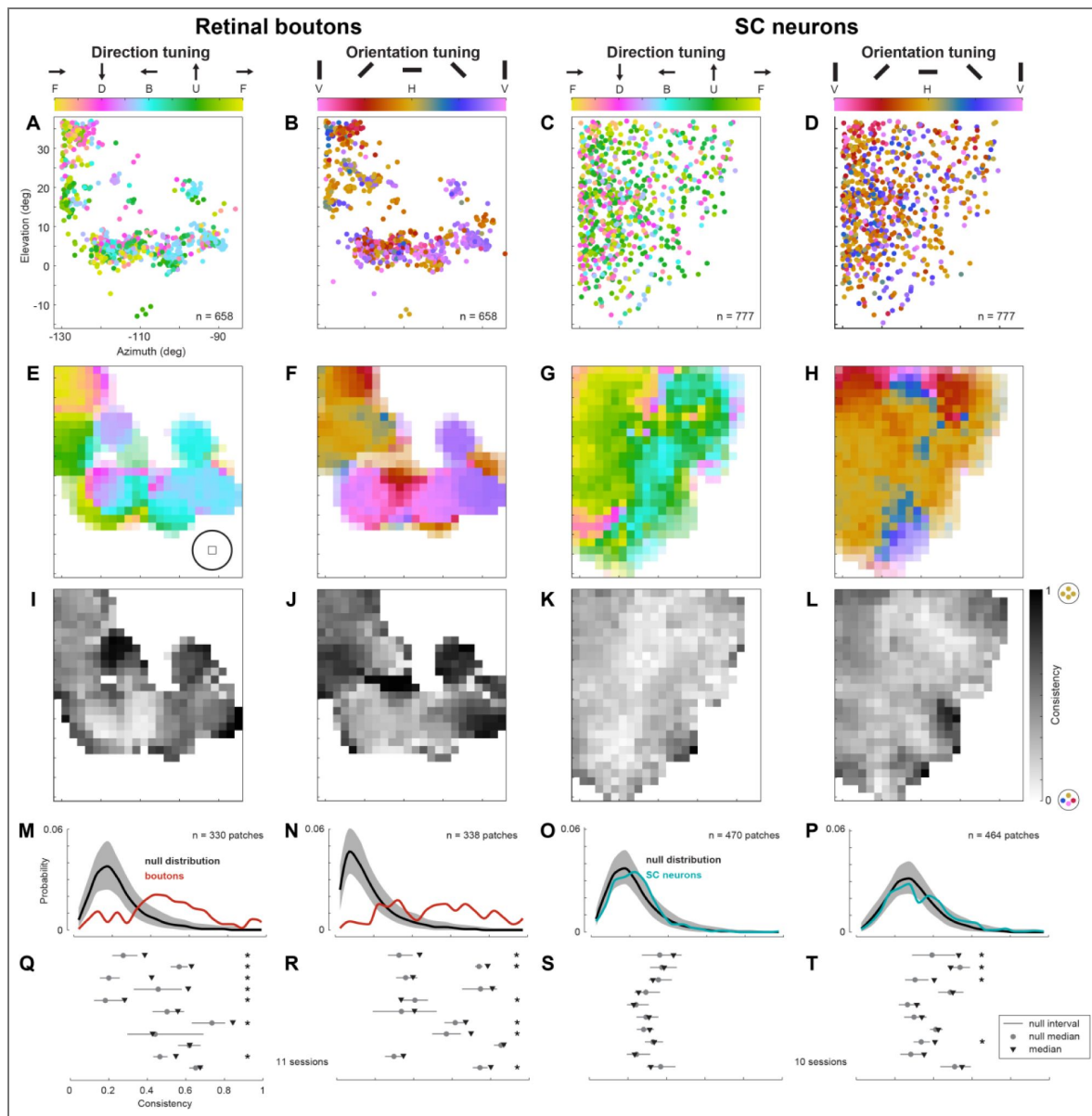


Figure 4. Local populations showed low consistency of direction and orientation preferences.

A–D. Direction (A) and orientation (B) preferences for all selective retinal boutons, plotted at their mapped RF locations in visual space. Corresponding data for SC neurons in C and D. For neurons, RF locations are based on the retinotopic fits (Figure 2). E–H. Mean direction (E) and orientation (F) preferences obtained by averaging across boutons within circular patches of 10° diameter in visual space (illustrated by circle in the lower-right corner). Transparency indicates patches with fewer than 20 boutons. Corresponding data for SC neurons in G and H. I–L. Consistency of direction (I) and orientation (J) preferences within 10° patches, quantified as the inverse circular variance of preferences (same pooling as in E and F) for retinal boutons. Corresponding data for SC neurons in K and L. M–P. Distributions of consistency for direction (M) and orientation (N) across patches (red; at least five units per patch) compared to null distribution obtained by permuting RF locations across boutons (black line: median; gray band: 2.5th–97.5th percentile interval). Corresponding data for SC neurons in O and P. RF locations for SC neurons are based on retinotopic fit (Figure 2). Q–T. Mean consistency in direction (Q) and orientation (R) preferences across all patches per recording for retinal boutons (black triangles), compared to corresponding null distribution for each recording (gray; dot: median; vertical line: 2.5th–97.5th percentile interval). Recordings with significant consistency values are marked by a star. Corresponding data for SC neurons in S and T.

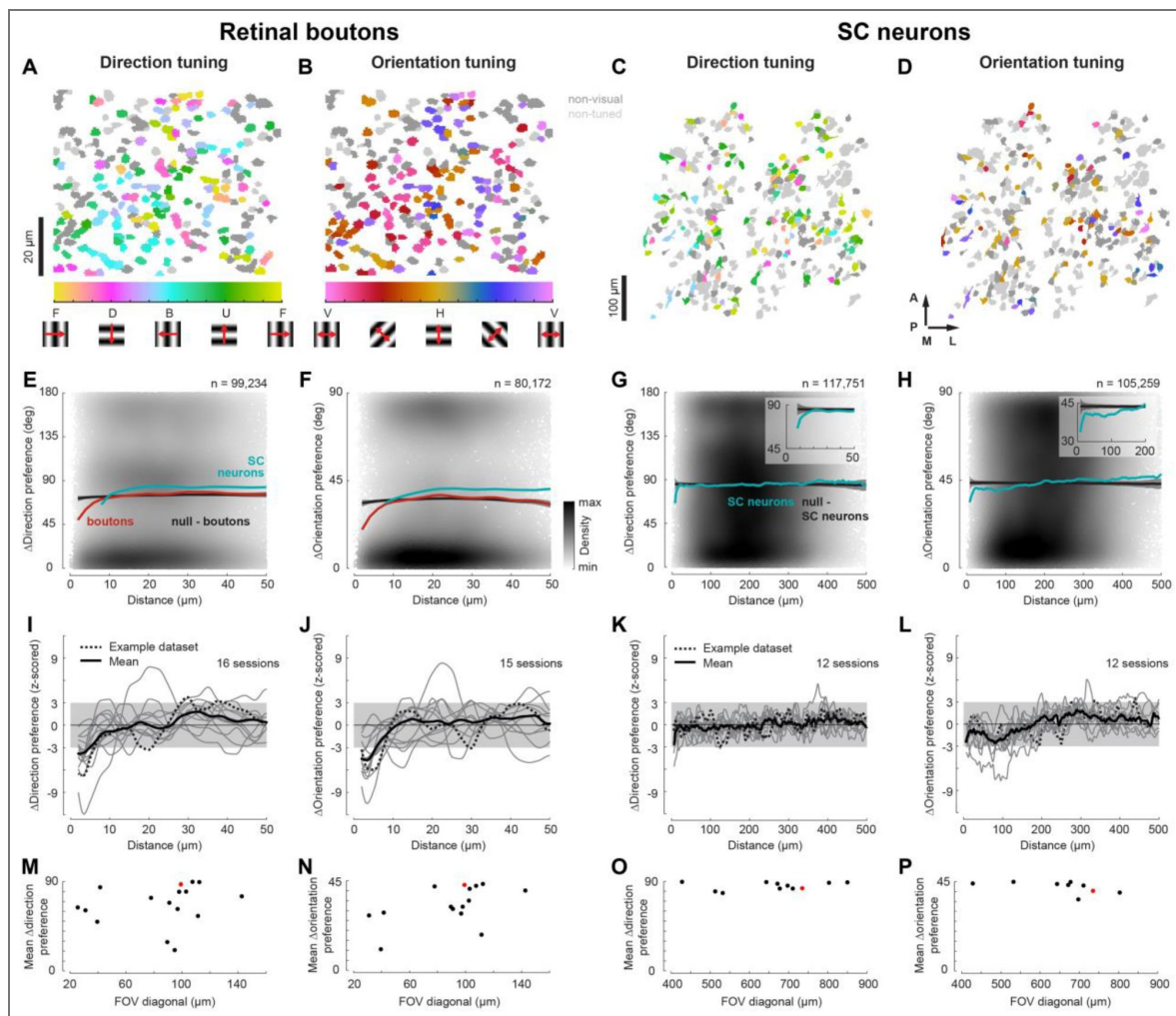


Figure 5. Pairwise similarity of direction and orientation tuning was largely independent of lateral distance.

A–D. Examples of direction (A) and orientation (B) preferences of individual retinal boutons in one FOV. Each bouton is plotted at its imaging location; non-responsive boutons are shown in dark gray and non-selective boutons in light gray. Corresponding data for SC neurons in C and D. Neurons from four imaging planes are shown. E–H. Pairwise difference in direction (E) and orientation (F) preferences as a function of lateral distance in the SC (depth ignored) for all pairs of simultaneously recorded retinal boutons (dots colored by density). Mean difference as a function of distance (red) is compared to the null distribution obtained by permuting bouton locations (black line: median; gray band: 2.5th–97.5th percentile interval). For comparison, mean pairwise difference of SC neurons is overlaid (teal; same data as in G, H). Corresponding data for SC neurons in G and H. Insets show zoom-in. I–L. Mean pairwise difference in direction (I) and orientation (J) preference for each recording (gray lines, at least 15 selective units each) as a function of distance, normalized to the null distribution of that recording. Mean (horizontal black line) and ± 3 SEMs (gray shade) of the null distribution are shown for comparison. Example recordings from A and B are shown as dotted lines; mean across all recordings depicted as bold black line. Corresponding data for SC neurons in K and L. M–P. Mean pairwise difference in direction (M) and orientation (N) preference for each recording (at least 15 selective units each) as a function of FOV size (length of diagonal). Example from A and B shown in red. Corresponding data for SC neurons in O and P.

boutons. This was not the case. Mean pairwise differences in bouton recordings showed no systematic relationship with FOV size, and some bouton recordings exhibited mean differences as large as those in neuron recordings with FOVs at least four times bigger (Figure 5M–P [↗](#)).

This weak clustering persisted when we only considered strongly selective units, when we measured distances in visual space rather than in FOVs, and when we used different visual stimuli. Restricting the analysis to DS-only or OS-only units (as previously by De Malmazet et al., 2024 [↗](#), 2018) did not reveal additional direction clustering and only modestly increased orientation similarity (Figure S5A–D [↗](#)). When we considered pairwise differences as a function of RF distance, bouton and neuron pairs showed the same pattern, with deviations from chance confined to RF pairs within a few degrees (Figure S5E–H [↗](#)). Pairwise tuning differences derived from moving bars or static gratings were similar to those for drifting gratings, with only SC neurons within ~20 μm showing clearly elevated similarity (Figure S5I–K [↗](#)).

Tuning across SC depth was heterogeneous and weakly related to the retinal topography

Neuropixels recordings across the full depth of the SC revealed robust tuning to motion direction and orientation. We recorded spiking activity using Neuropixels probes inserted vertically through the full depth of the SC in an experimental setup similar to the two-photon imaging experiments (Figure 6A [↗](#), B [↗](#)). Single units were isolated by automatic spike sorting using Kilosort2.5 (Pachitariu et al., 2024 [↗](#)) and quality metrics followed by manual curation. The border between sSC and deep SC (dSC) was determined from visually evoked local field potentials and current source density (Figure S6A–C [↗](#)). Around 48% of neurons in the SC were well tuned (Figure 6C–E [↗](#), I [↗](#), J [↗](#), K [↗](#)) and the population was biased towards cardinal directions and orientations (Figure 6F [↗](#), G [↗](#)). Preferred directions were strongly correlated with preferred orientations (Figure S6D [↗](#)). Direction selectivity in sSC was generally higher than orientation selectivity (Figure 6K [↗](#)), while, across all depths, OS units were more numerous than DS units (Figure S6E [↗](#)).

Direction and orientation preferences showed no consistent pattern with SC depth. When sorting preferred directions and orientations by depth within each recording no systematic trend or layering was revealed (Figure 6H [↗](#), J [↗](#)), even though RFs of simultaneously recorded neurons largely overlapped (Figure 7A [↗](#), B [↗](#)). Quantitatively, median pairwise tuning differences were close to the values expected for a uniform distribution of preferences (90° for direction, 45° for orientation) and rarely deviated from the medians obtained after randomly permuting units across recordings (Figure 6L [↗](#), N [↗](#)). Pairwise differences were also independent of distance in depth between neurons: mean tuning differences as a function of depth separation closely followed null distributions based on permuted depths for both direction and orientation (Figure 6M [↗](#), O [↗](#)).

Direction and orientation tuning across the depth of SC showed little resemblance to the retinal spherical topography. As before, we located RFs of recorded neurons within the visual field (Figure 7A–C [↗](#)). When we compared SC tuning to the geometric model derived from retinal RGCs (Laniado et al., 2025 [↗](#); Sabbah et al., 2017 [↗](#)), the match between measured and predicted preferences in sSC was weaker than in retinal boutons (Figure 3 [↗](#)) and weakest for dSC neurons (Figure 7D–M [↗](#)). In particular, direction and orientation preferences of dSC neurons showed only marginal alignment with the predicted retinal vectors, which was as weak as expected by uniform distributions of preferences (Figure 7L [↗](#), M [↗](#)). Together, these results are consistent with increasing integration across diverse retinal inputs as one moves away from the retinal input. We noticed that the distributions of direction and orientation classes recorded in sSC using electrophysiology (Figure 7F [↗](#), H [↗](#)) appear to be different from the distributions in imaged sSC neurons (Figure 3J [↗](#), K [↗](#)). This may be explained by the difference in RF locations, differences in recorded depth (imaging is biased to the most superficial sSC), or individual differences across mice.

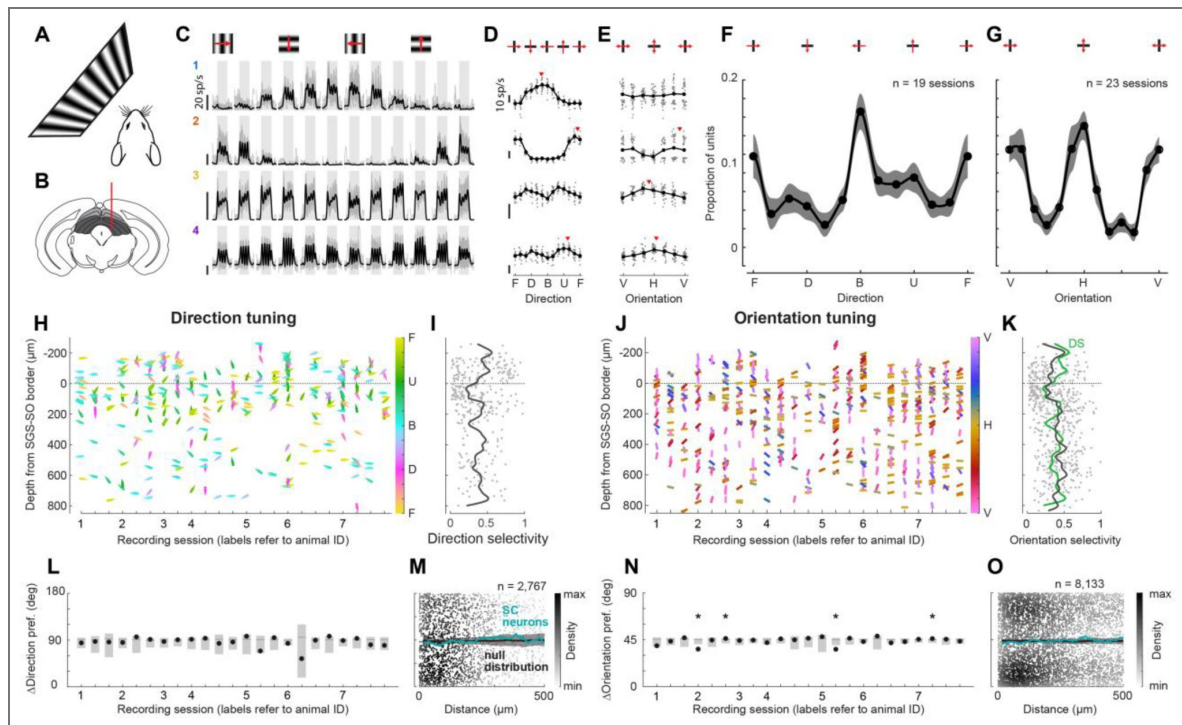


Figure 6. Electrophysiological recordings across SC depth confirmed the lack of functional clustering.

A. Visual stimuli were presented on a single monitor covering -95° to 0° azimuth and -27° to 51° elevation. Drifting grating stimuli covered the full screen. B. Neuropixels probes were used to record single units from sSC (light gray) and dSC (dark gray). C. Single-trial (gray) and mean (black) spike rate traces of example units in response to sinusoidal gratings drifting in 12 directions. Grating presentations (2 s each) are indicated by gray shading. D, E. Direction (D) and orientation (E) tuning curves for example units in C, showing single-trial response amplitudes (gray dots), mean amplitudes (black dots), and preferred directions and orientations if significantly tuned (red triangles). F, G. Mean distribution ($\pm$ SEM) of preferred directions (F) and orientations (G) across all recording sessions. Datasets with fewer than 10 selective units were excluded. H. Preferred direction of all recorded units, as a function of depth in SC for each recording session (multiple sessions per animal). I. Direction selectivity of all units as a function of depth in SC. J, K. Same as in H and I, for orientation tuning. For comparison, direction selectivity profile (green) from I is overlaid in K. L. Mean of pairwise direction difference of each recording (dots), compared to corresponding null distribution (gray line: median; light-gray box: 2.5th–97.5th percentile interval). M. Pairwise difference in direction preference as a function of depth separation in SC for all pairs of simultaneously recorded units (dots colored by density). Mean difference as a function of separation (teal) is compared to null distribution obtained by permuting unit depths (black line: median; gray band: 2.5th–97.5th percentile interval). N, O. Same as in L and M, for orientation tuning.

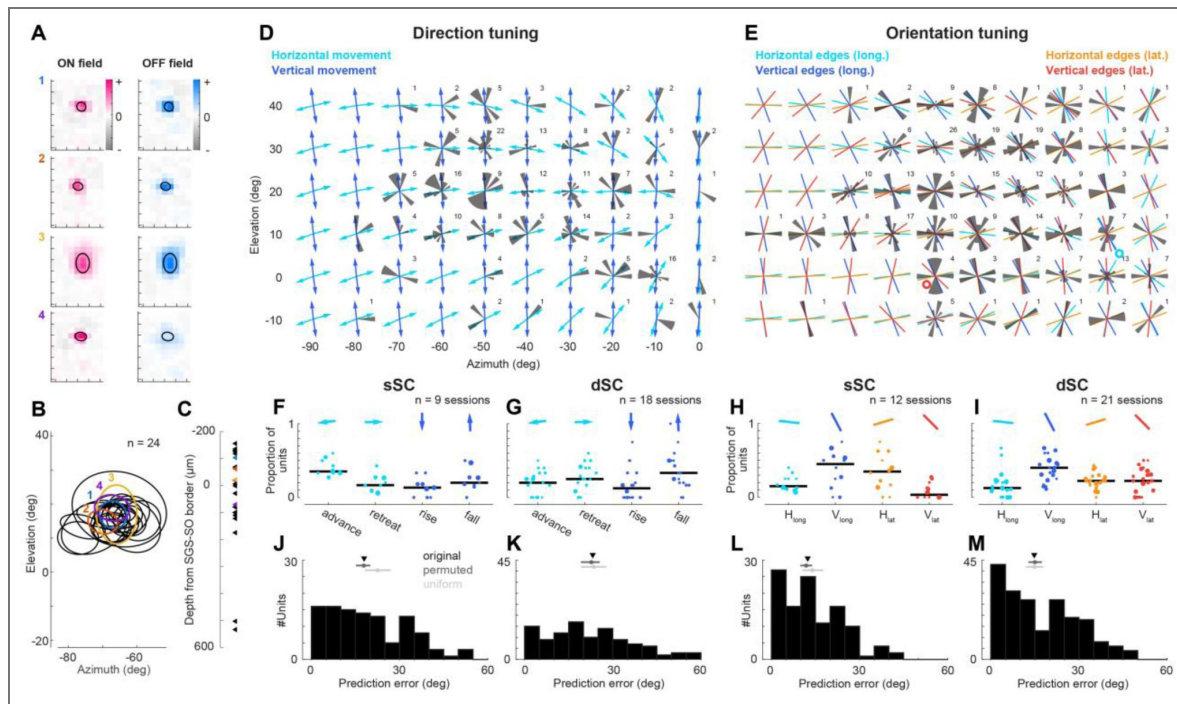


Figure 7. Retinal topography was progressively weakened from superficial to deep SC.

A. ON and OFF RFs of four example units (same as in Figure 6C). RF outlines (center ± 1 SD of fitted 2D Gaussian) shown in black. B. RF outlines of all units recorded in one session, including examples in A (colored). C. Depth distribution of units with RFs from B. Depths were aligned to the border between sSC and dSC. D, E. Direction (D) and orientation (E) preferences of SC units across visual space, plotted at their mapped RF locations (not retinotopic). Arrows in D point along directions defined by DSGC longitudes (see Figure 3A). Colored lines in E indicate orientations of longitudinal and latitudinal fields for OSGCs (see Figure 3B, C). Numbers indicate population size in each polar histogram. The depicted region of visual space in E includes two poles of the spherical axes: one for horizontal longitudes (cyan circle) and one for vertical latitudes (red circle). F–I. Classification of all DS units in sSC (F) and dSC (G) according to closest predicted direction vector. Each dot shows one recording session (dot size reflects total number of units); black line indicates median proportion across sessions. Arrows at the top illustrate example predicted vectors from a central location in D. Corresponding data for OS units in H and I. J–M. Angular differences between measured direction preferences and the closest predicted vectors in sSC (J) and dSC (K). The median difference (triangle) is compared to the distribution obtained by permuting RF locations (dark gray: median; band: 2.5th–97.5th percentile interval) and to a uniform preference model (light gray: median; band: 2.5th–97.5th percentile interval). sSC: 18.0° original, 15.2°–20.0° permuted, 18.3°–27.0° uniform; dSC: 23.2° original, 19.2°–25.6° permuted, 19.2°–27.9° uniform. Corresponding data for OS units in L and M. sSC: 12.4° original, 9.9°–13.9° permuted, 10.7°–17.7° uniform; dSC: 14.8° original, 12.8°–17.4° permuted, 12.0°–17.9° uniform.

Discussion

Our results demonstrate that the topographic organization of direction and orientation tuning in mouse SC was largely inherited from the retina and then transformed within SC. Using two-photon calcium imaging of retinal boutons and SC neurons in posterior monocular SC, together with Neuropixels recordings across the depth of anterior binocular SC, we found robust tuning and shared cardinal biases in both retinal inputs and neurons. Retinal boutons closely followed retinal geometric models that predict direction and orientation preferences in retina-centered coordinates (Laniado et al., 2025 [DOI](#); Sabbah et al., 2017 [DOI](#)), while the alignment to the geometric model decreased from retinal boutons to superficial neurons and further to deep neurons. Local clustering of tuning was weak in all neural populations and confined to very small spatial scales (10–20 μm), while tuning preferences across visual space lacked smooth gradients and stereotyped patterns.

Maps or no maps?

Pairwise differences in preferred direction between SC neurons in our data were largely independent of lateral distance, and consistency of preferences within 100- μm neighborhoods was at chance. These findings agree with some studies (Chen et al., 2021 [DOI](#); Inayat et al., 2015 [DOI](#)), but disagree with reports of local clustering of either direction or orientation preferences (Ahmadlou and Heimel, 2015 [DOI](#); De Malmazet et al., 2018 [DOI](#); Feinberg and Meister, 2014 [DOI](#); Kasai and Isa, 2021 [DOI](#); Li et al., 2020 [DOI](#)). But among cluster- or map-positive studies, the strength of pairwise differences varied markedly: for pairs <100 μm apart, median direction differences were $\sim 40^\circ$ in some studies (De Malmazet et al., 2018 [DOI](#); Li et al., 2020 [DOI](#)) versus $\sim 80^\circ$ in others (Kasai and Isa, 2021 [DOI](#)), and median orientation differences ranged from $\sim 30^\circ$ (Feinberg and Meister, 2014 [DOI](#); Kasai and Isa, 2021 [DOI](#)) to $<10^\circ$ (De Malmazet et al., 2018 [DOI](#)). At the level of retinal inputs, nearby boutons were reported to have almost identical orientation preferences and more homogeneous direction tuning than in our data (De Malmazet et al., 2024 [DOI](#)), whereas we observed clear clustering only within $\sim 10 \mu\text{m}$, plausibly reflecting multiple boutons from single RGCs (Hong et al., 2011 [DOI](#)).

Inconsistencies among previous reports are even more pronounced at the level of global maps. For direction, different studies emphasized a predominance of upward motion, upward-anterior motion, or opposing posterior/anterior biases in monocular versus binocular SC (Ahmadlou and Heimel, 2015 [DOI](#); De Malmazet et al., 2018 [DOI](#); Dräger and Hubel, 1975 [DOI](#); Li et al., 2020 [DOI](#)). For orientation, neurons near the center of the visual field were described as horizontally biased or largely non-selective (Ahmadlou and Heimel, 2015 [DOI](#); De Malmazet et al., 2018 [DOI](#)). Proposed map layouts range from concentric orientation maps centered on the nose to $\sim 30^\circ$ -wide patches of similar orientation without global structure (Ahmadlou and Heimel, 2015 [DOI](#); De Malmazet et al., 2018 [DOI](#); Feinberg and Meister, 2014 [DOI](#)), and large differences across individual animals are evident within single studies (e.g., Figure S3 [DOI](#) in De Malmazet et al., 2018 [DOI](#); Figure 6C [DOI](#) in Li et al., 2020 [DOI](#)). Against this heterogeneous background, our observation of broadly distributed, weakly clustered preferences suggests that strong, stereotyped maps are unlikely to be a universal organizing principle of SC.

Several factors may contribute to the discrepancies across studies. Some focused on anterior SC and the binocular/monocular border, others on more posterior monocular regions, yet both map-like and salt-and-pepper organizations have been reported in both zones, so retinotopic location alone cannot explain the differences. Visual stimuli varied widely (drifting or static gratings, moving bars, dots), distortions from presenting spherical coordinates on flat screens were not always corrected, and some RF estimates were based on response delays of DS units to moving bars, which can misplace RFs and thus inferred maps (De Malmazet et al., 2024 [DOI](#), 2018). Most experiments used awake animals, but internal state mainly modulates gain rather than preferred direction or orientation (Chen et al., 2021 [DOI](#); Ito et al., 2017 [DOI](#); Savier et al., 2019 [DOI](#); Schröder et al.,

2020), making it an unlikely source of disagreement. Finally, biases in cell-type sampling are plausible if stimuli drive only a minority of neurons, implying that different functional subpopulations were emphasized.

A partial resolution emerges when considering the four cardinal direction and orientation preferences that are predicted by the retinal topographic organization to vary smoothly across visual space (Laniado et al., 2025; Sabbah et al., 2017). This framework encompasses concentric orientation layouts near the front, in addition to other geometries. Systematic or random shifts in the relative strengths of these cardinal channels across animals, recording methods, or stimulus sets could change which preference appears dominant and produce various apparent maps. Moreover, some studies analyzed DS and OS populations separately and reported strong structure within, but random mixing between, these groups (De Malmazet et al., 2024, 2018). Given that neurons with both direction and orientation selectivity typically exhibit approximately orthogonal preferences, pooling DS and OS populations and relaxing selectivity criteria will naturally yield more heterogeneous local preference distributions.

Retinal topography was progressively transformed by collicular circuits

Retinal boutons in sSC closely followed the topographic models of DSGCs and OSGCs (Laniado et al., 2025; Sabbah et al., 2017; Tiriach et al., 2022). This is expected because 85–90% of RGCs, including all four cardinal DSGC types, project to SC (Ellis et al., 2016; Kay et al., 2011), their terminals predominantly target the superficial layers that we imaged (Dhande and Huberman, 2014; Hong et al., 2011; Huberman et al., 2009), and their projections follow strict retinotopy (Molotkov et al., 2023; Sibille et al., 2022). Our RF measurements confirmed this tight retinotopic alignment.

In sSC neurons, the topographic organization remained evident for direction but was weaker for orientation. Direction selectivity and preferred direction in SC are largely inherited from DSGCs and amplified by similarly tuned intracollicular inputs (Shi et al., 2017; Sibille et al., 2022), while some DS neurons in SC also receive input from non-DSGCs (Kay and Triplett, 2017). Axis-selective responses may be inherited from OSGCs, or may arise from convergent innervation by oppositely tuned DSGCs or local SC circuits (Kay and Triplett, 2017). Our finding that direction preferences of superficial neurons remained well aligned with the retinal topography, whereas orientation preferences deviated more than in boutons, is therefore consistent with a largely preserved feedforward scaffold for direction combined with mixing of multiple retinal channels and local circuitry that blurs topographic orientation organization at the single-neuron level. The weaker agreement with the topographic organization in Neuropixels-recorded superficial neurons likely reflects undersampling of the most superficial sSC layer, which is preferentially targeted by DSGCs.

In dSC, we found almost no agreement with the retinal topographic organization despite robust tuning. Functional classifications in previous studies identified “RGC-like” neuron types that predominate in sSC and more “SC-specific” types that become more prominent with depth (Lee et al., 2020; Li and Meister, 2023), consistent with increasing transformation away from retinal channels. This transformation in dSC is expected given further functional integration within SC and the convergence of signals from other sources, including cortical and brainstem structures (Cang et al., 2024; Liu et al., 2022; Wheatcroft et al., 2022). Together, these results support a hierarchical view in which a retinally derived topographic organization is preserved in retinal boutons, but increasingly mixed across SC layers.

Parallels with mouse V1 and theoretical constraints on map organization

The functional organization we observed in mouse SC closely resembles that in mouse V1. In both areas, neurons are arranged in precise retinotopic maps and are robustly tuned to orientation and direction, yet their preferences form a largely salt-and-pepper pattern with weak local clustering,

rather than classical orientation columns (Bonin et al., 2011 [↗](#); Chen et al., 2021 [↗](#); Inayat et al., 2015 [↗](#); Kaschube, 2014 [↗](#); Niell and Stryker, 2008 [↗](#)). V1 shows short-range clustering on the scale of tens to hundreds of micrometers for orientation and, more modestly, for direction (Kondo et al., 2016 [↗](#); Ringach et al., 2016 [↗](#); Yu et al., 2025 [↗](#)), comparable to the very small-scale clustering we found in SC. Local orientation biases in V1 have been linked to ON/OFF RF tiling in the retina (Jimenez et al., 2018 [↗](#)), whereas SC may achieve better local coverage by directly inheriting direction tuning from DSGCs and orientation tuning from OSGCs or from pooling oppositely tuned DSGCs.

Theoretical accounts of map formation suggest that such salt-and-pepper organization is expected in mouse SC. Across species, rodents and rabbits exhibit salt-and-pepper distributions of tuning, whereas cats, ferrets, and primates display columnar maps (Kaschube, 2014 [↗](#)), though not for all visual features (Martin and Schröder, 2013 [↗](#); Yen et al., 2007 [↗](#)). Models based on the ratio of retinal inputs to cortical area—the mapping or sampling ratio—predict that low ratios favor a salt-and-pepper organization, whereas high ratios favor columnar maps (Ibbotson and Jung, 2020 [↗](#); Jang et al., 2020 [↗](#); Najafian et al., 2022 [↗](#); Schmidt and Wolf, 2021 [↗](#)). Published estimates place SC in a low-mapping-ratio regime: a larger fraction of RGCs projects to SC than to the thalamocortical pathway (Ellis et al., 2016 [↗](#); Martin, 1986 [↗](#)), SC RFs are smaller than those in V1 (Niell and Stryker, 2008 [↗](#); Wang et al., 2010 [↗](#)), and SC has a smaller surface area (Edwards et al., 1986 [↗](#); Mazade and Alonso, 2017 [↗](#)). The same anatomical constraints that favor salt-and-pepper organization in mouse V1 therefore naturally extend to SC.

Functional implications and future directions

Biases toward directions aligned with body and gravity axes likely facilitate decoding of self-motion and ethologically relevant motion (Sabbah et al., 2017 [↗](#); Tiriuc et al., 2022 [↗](#)), whereas local salt-and-pepper organization ensures that SC can represent any motion direction or edge at each location and support diverse behaviors (Hoy and Farrow, 2025 [↗](#); Wheatcroft et al., 2022 [↗](#)). The spherical organization of OSGCs has been proposed to make it easy for a downstream decoder to infer edge orientation from the relative activity of a few OSGC types (Laniado et al., 2025 [↗](#)). In the binocular zone, the vertical cell type of the longitudinal system and the horizontal type of the latitudinal system provide the highest decoding efficiency (Laniado et al., 2025 [↗](#)). These two classes were the most common among sSC neurons recorded with Neuropixels in the binocular zone (Figure 7 [↗](#)). This convergence indicates that SC receives particularly informative retinal orientation channels, even though their expression at the neuronal level is weakly clustered.

Future work should extend these measurements to larger SC populations while preserving circuit integrity. Ideally, this would combine broad retinotopic coverage, including anterior/binocular and posterior/monocular SC, with single-cell resolution. Optical methods are attractive but usually require removal of overlying cortex, which risks altering cortical input or damaging sSC; deep imaging approaches that access SC without cortical removal would therefore be especially valuable. New experiments should also quantify variability across individuals, given the substantial inter-animal differences reported previously, and test how organization generalizes to richer stimulus conditions. Natural images and movies, motion-contrast stimuli, and size- or context-dependent motion could reveal structure not engaged by simple gratings and bars (DePiero et al., 2024 [↗](#)). Finally, relating SC organization to behavior—for example, by testing whether directions and orientations near overrepresented preferences are discriminated more accurately, or whether mice move head or eyes to bring stimuli into specific visual-field regions—will be useful to link the retinal scaffold and its collicular transformation to concrete behavioral advantages.

Methods

Experimental Model and Subject Details

All procedures were conducted in accordance with the UK Animals Scientific Procedures Act (1986) under personal and project licenses released by the Home Office following appropriate ethics review.

For two-photon imaging, we used 1 inbred C57Bl/6J (www.jax.org/strain/000664; 1 male) and 12 mice (7 females, 5 males) obtained by crossing Gad2-IRES-Cre (www.jax.org/strain/010802) and Ai9 (www.jax.org/strain/007909). The heterozygous offspring expressed TdTomato in glutamate decarboxylase 2-positive (GAD2+) cells to identify inhibitory neurons. For electrophysiological recordings, we used 4 Vgat-IRES-Cre mice (www.jax.org/strain/016962; 2 females, 2 males) and mice (1 female, 2 males) obtained by crossing Vgat-IRES-Cre (www.jax.org/strain/016962) and Ai14 (www.jax.org/strain/007908). Animals were 9–29 weeks old at the time of surgery and were used for experiments up to the age of 54 weeks. Mice were kept on a 12-h light: 12-h dark cycle. Most animals were single housed after the first surgery.

Overview of collected data

	Retinal boutons (2P)	SC neurons (2P)	SC neurons (ephys)
Animals	8	5	7
Recording sessions	17	12	23
Recorded units	3,677	4,769	1,474
Units responsive to drifting gratings	2,391	2,378	1,120
Selective units	523 DS-only; 322 OS-only; 1,056 DS & OS	332 DS-only; 213 OS-only; 1,243 DS & OS	128 DS-only; 373 OS-only; 211 DS & OS
Units for which RFs could be mapped	1,979 of 3,282 recorded units (13 sessions, 8 animals)	2,471 of 4,769 recorded units (12 sessions, 5 animals)	539 of 1,120 recorded units (23 sessions, 7 animals)
Units with RF and tuned to direction or orientation	1,084 (of 1,274 DS); 1,056 (of 1,226 OS)	1,055 (of 1,575 DS); 955 (of 1,456 OS)	224 (of 339 DS); 308 (of 584 OS)

Surgical procedures

Surgical procedures for animals undergoing two-photon imaging were described previously (Schröder et al., 2020). Here, we provide a shorter description and point out differences to animals undergoing electrophysiological experiments. Animals were anesthetized with isoflurane and the following drugs were administered subcutaneously: carprofen (5 mg/kg; Rimadyl, Pfizer) and dexamethasone (0.5 mg/kg; Colvasone, Norbrook) in animals undergoing surgery for two-photon imaging experiments (2P); or meloxicam (5 mg/kg; Metacam, Boehringer Ingelheim), buprenorphine (0.05 mg/kg; Vetergesic, Ceva Animal Health), and dexamethasone (4 mg/kg; Rapidexon, Dechra) in animals undergoing surgery for electrophysiology experiments (ephys). The scalp was shaved and disinfected, and local analgesia (2P: lidocaine: 6 mg/kg, Hameln pharmaceuticals ltd; ephys: bupivacaine: 7 mg/kg, Marcain, AstraZeneca) was injected subcutaneously under the scalp prior to the incision. Eyes were covered with eye-protective gel (Chloramphenicol, Martindale Pharmaceuticals Ltd). A local anesthetic (2P: 5% Lidocaine ointment, TEVA UK; ephys: 5% EMLA cream, Aspen) was applied to the ear bars, before the animal was placed into a stereotaxic apparatus. The skin covering and surrounding the area of interest was removed, and the skull was cleaned of connective tissue. A custom made headplate was positioned above the area of interest and attached to the bone with Superbond C&B (Sun Medical). Throughout all surgical procedures, the animal was kept on a heating pad to stabilize body

temperature at 37°C. Subcutaneous injections of 0.01 ml/g/h of Sodium Lactate (Hartmann's solution) were given. After the surgery, the animal was placed into a cage on a heating pad (37°C) for recovery from anesthesia. Mice were given three days to recover while being treated with Carprofen or Metacam.

To image retinal boutons in SC, we cloned a variant of GCaMP6f fused with a localization signal targeting synaptic terminals (SyGCaMP6f) and packaged it into an AAV2/2 vector (Addgene, 51085) so that GCaMP expression was restricted to boutons (Dreosti et al., 2009 [DOI](#)). We injected approximately 3 µl of AAV2/2.hSyn1.SyGCaMP6f.SV40 with a concentration of 2.44×10^{12} viral particles/ml into the vitreous humor of the left eye. In all animals used for two-photon imaging, a circular 4 mm craniotomy (centered at approximately -4.2 mm AP and 0.5 mm ML from Bregma) was made. For two-photon imaging of activity in SC neurons, we injected the virus AAV2/1.Syn.GCaMP6f.WPRE.SV40 (Chen et al., 2013 [DOI](#)) at a final concentration of $2.30\text{--}4.39 \times 10^{12}$ GC/ml, at a rate of 2.3 nl every 6 s (Nanoject II, Drummond), below the surface of the right SC. As the posterior SC is covered by a large sinus running through the dura, we implanted a custom-made implant permanently to push the sinus anteriorly and gain optical access to the SC. The implant was fixed to the skull with Vetbond (3M) and Superbond C&B (Sun Medical).

For electrophysiological recordings, a craniotomy (centered at -3.7 mm AP and 0.6 mm ML relative to Bregma, in the right hemisphere) was performed above the anterior SC at least one day before the first acute recording was performed.

Two-photon imaging

Most of the two-photon imaging datasets have been used in a previous study (Schröder et al., 2020 [DOI](#)), and additional datasets have been collected in the same way. Shortly, two-photon imaging was performed using a standard resonant microscope (B-Scope, ThorLabs Inc.) equipped with a 16x, 0.8 NA water immersion objective (N16XLWD-PF, Nikon) and controlled by ScanImage 4.2 (Pologruto et al., 2003 [DOI](#)). Excitation light at 970–980 nm was delivered by a femtosecond laser (Chameleon Ultra II, Coherent). Multi-plane imaging was performed using a piezo focusing device (P-725.4CA PIFOC, Physik Instrumente, 400 µm range). Laser power was depth-adjusted and synchronized with piezo position using an electro-optical modulator (M350-80LA, Conoptics Inc.). Emission light was collected using two separate channels, one for green fluorescence (525/50 nm emission filter) capturing the calcium transients and one for red fluorescence (607/70 nm emission filter).

For imaging neurons in SC, we used 3–4 imaging planes separated by 9–30 µm at depths of 15–100 µm from the surface of SC. The field of view spanned 340–640 µm in both directions at a resolution of 512 x 512 pixels. The frame rate per plane was 6.0–7.5 Hz.

For imaging retinal boutons in SC we used 5 or 10 imaging planes (with 3 or 6 fly-back planes) with an inter-plane distance of <2 µm at depths of 8–46 µm from the surface of SC. The field of view spanned 42–135 µm and each plane was imaged at a rate of 7.5 Hz. Because the calcium indicator was localized to synaptic boutons, fluorescence was not affected by fibers of passage.

Electrophysiology

Recordings were made using Neuropixels electrode arrays (Jun et al., 2017 [DOI](#)). Probes were mounted to a custom 3D-printed holder attached to a steel rod and positioned with a micromanipulator (uMP-4, Sensapex Inc.). For subsequent track reconstruction, probes were coated with a dye (DiI: ThermoFisher Vybrant V22888/V22885; or DiO: ThermoFisher Vybrant V22886) by immersing the shank for 5–10 min in an Eppendorf tube containing the dye solution, after which they were removed and air-dried. Probes had a soldered connection between external reference and ground; at the headstage, ground was connected to an Ag/AgCl wire placed on the skull. Craniotomies and the ground wire were kept covered with saline throughout the experiment. Probes were advanced through the dura and lowered to the target depth at ~10 µm/s, then allowed to settle for 10 min before recording. Signals were acquired with SpikeGLX (<https://billkarsh.github.io/SpikeGLX> [DOI](#)), and were separated into an action-potential (AP) band

(hardware high-pass filtered at 300 Hz, sampled at 30 kHz) and a local-field-potential (LFP) band (low-pass filtered at 500 Hz, sampled at 2.5 kHz). Recordings were repeated across different penetration sites on multiple subsequent days.

Experimental setup and visual stimuli

The mouse was head fixed with a headplate holder that did not obstruct the visual field. For two-photon imaging, the mouse was free to run on an air-suspended Styrofoam ball (20 cm in diameter), whose rotation was measured by two optical computer mice (Dombeck et al., 2010). For electrophysiology, the mouse was free to run on a plastic wheel (8.5 cm wide, 19 cm diameter) (Warren et al., 2021 [DOI](#)), whose rotation was measured by a rotary encoder (1,024 pulses per rotation, Kübler, Germany). The mouse was acclimated to head-fixation for at least three days before the first recording session, step-wise increasing fixation time from ≥ 5 min on the first day to ≥ 20 min on the last day.

For two-photon imaging, the mouse was surrounded by three computer screens (Iiyama ProLite E1980SD placed ~ 20 cm from the mouse's eyes; 60 Hz refresh rate) at right angles covering the visual field from -135° to 135° azimuth and -42° (bottom) to 42° (top) elevation. In some experiments, Fresnel lenses (BHPA220-2-6 or BHPA220-2-5, Wuxi Bohai Optics) were mounted in front of the monitors to compensate for reduction in luminance and contrast at steeper viewing angles relative to the monitors, and lenses were coated with scattering window film (frostbite, The Window Film Company) to prevent specular reflections. The red channel of the monitors was switched off to reduce light contamination in the red fluorescence channel.

For electrophysiological recordings, visual stimuli were presented on one computer screen (Dell P1917S; 37.5×30 cm, 1280×1024 pixels, 60 Hz refresh rate), positioned 20 cm from the mouse's left eye and spanning -95° (left) to 0° (right) azimuth and -27° (bottom) to 51° (top) elevation.

Brightness levels of the monitors were linearized around a mean level of 83 cd/m^2 (two-photon imaging) or 36 cd/m^2 (electrophysiology). All visual stimuli, whose sizes and positions were defined with respect to the visual field, were mapped onto the viewing plane(s) of the computer screen(s) and were created using PsychToolbox (Brainard, 1997 [DOI](#)) or BonVision (Lopes et al., 2021 [DOI](#)).

Sinusoidal drifting gratings covered the complete screen space (full-field), had a spatial frequency of 0.08 cycles/deg, a temporal frequency of 2 cycles/s, and were presented at 100% contrast. Gratings drifted in one of 12 equidistant directions and were presented for 2 s separated by a gray screen for 3–6 s (two-photon imaging) or 2–3 s (electrophysiology).

Static sinusoidal gratings also covered the complete screen space, had a spatial frequency of 0.08 cycles/deg, and were presented at 100% contrast. Static gratings varied in orientation (0° – 135° , in steps of 45°) and spatial phase (0° , 120° , 240°), comprising all possible combinations. Each static grating was presented for 1 s with an inter-stimulus interval of 2–3 s.

Moving bars were black on mean-gray background, had a width of 2.5° , a length spanning the full screen size, and moved at a speed of $15^\circ/\text{s}$. Bars varied in direction of movement (0° – 315° , in steps of 45°). Each bar was presented in a window of 4 s with an inter-stimulus interval 3–5.5 s.

Within blocks of the same stimulus type, each stimulus was repeated 15 or 20 times following a pseudo-random presentation order.

To map RFs and to determine the position of SC along the probe, we presented checkerboard images (visual noise) with white, black and mean gray squares with an edge length of 10° . The stimulus updated at a rate of 6 Hz and was presented for >9 min. In each noise image, each square was randomly assigned its luminance value with a 98% probability of being gray and 1% probabilities of being black and white (for two-photon imaging), or 20 out of 288 squares was black or white (for electrophysiology). Alternatively, RFs were mapped using black and white circles with diameters varying between 0.5° and 32° . Circles were presented individually on gray background at a rate of 10 Hz, for at least 5 min.

Data analysis

Preprocessing of two-photon imaging data

Preprocessing has been described previously (Schröder et al., 2020 [DOI](#)). Shortly, all raw two-photon imaging movies were analyzed using Suite2p (implemented in Matlab, Mathworks) to align frames and detect regions of interest (Pachitariu et al., 2016 [DOI](#)). We used the red channel reflecting TdTomato expressed in inhibitory neurons to align frames. For imaging data of retinal boutons, we also aligned frames in depth, i.e. when the brain moved perpendicular to the imaging planes, fluorescence data from neighboring imaging planes was used to correct this movement. Regions of interest (ROIs) were detected using the aligned frames of the green channel, and were then manually curated using the Suite2p GUI.

Using the aligned movies and detected ROIs resulting from Suite2p analysis, we extracted the fluorescence from the green and the red channel within each ROI. To correct the calcium traces for contamination by surrounding neuropil, we also extracted the fluorescence of the surrounding neuropil for each ROI using the green channel. To correct for contamination, the neuropil trace, N , was subtracted from the calcium trace, F , using a correction factor α : $F_c(t) = F(t) - \alpha \cdot N(t)$. The correction factor was determined for each ROI individually (Dipoppa et al., 2018 [DOI](#)). To correct for potential brain movements, we used the red fluorescence traces of each ROI to regress out changes in fluorescence that were not due to neural activity.

To avoid sampling the same neuron/bouton more than once, we removed duplicate ROIs that were close to each other in neighboring imaging planes and that had highly correlated calcium traces.

Spike sorting

Preprocessing and spike sorting were performed using the IBL-sorter pipeline (International Brain Laboratory et al., 2022 [DOI](#)), which includes common-median referencing, channel-noise whitening, detection and removal of noisy channels, and an initial estimate of probe drift. Kilosort2.5 was then executed using the detection thresholds of 9 for high-threshold detection and 3 for low-threshold detection. Unit quality was assessed with Bombcell (Fabre et al., 2023 [DOI](#)). Units were included only if they were labelled *good* by Bombcell and satisfied the following quality criteria: refractory-period violations $\leq 15\%$, $< 35\%$ of spikes falling below the detection threshold, presence ratio $> 25\%$, and ≥ 300 spikes. All retained units were subsequently curated manually in phy (<https://github.com/cortex-lab/phy> [DOI](#)) to identify potential merges or splits. Decisions were based on waveform similarity, the bimodality of waveform feature distributions, the spike amplitude distributions and drift patterns, and cross-correlogram structure between putative units.

Position of SC along recording probe

To identify the SC surface and the border between stratum griseum superficiale (SGS) and stratum opticum (SO) in Neuropixels recordings, we used visually evoked local field potentials (LFPs) following established approaches (Ito et al., 2017 [DOI](#); Lee et al., 2020 [DOI](#); Stitt et al., 2013 [DOI](#); Zhao et al., 2014 [DOI](#)). For each recording, we first subtracted the median LFP across time from each channel to remove channel-specific offsets. We then processed the two vertical columns of Neuropixels channels separately. Within each column, we smoothed the raw LFP in time and depth by convolving with a Gaussian kernel (5 channels in depth, 21 time bins). At each time point, we subtracted the LFP averaged across all channels of a column to emphasize local depth-dependent structure. We next identified the most effective pixel (for noise stimuli) or circle (for circle stimuli) that resulted in the largest LFP amplitude and aligned the LFP to the pixel's or circle's onset times.

The SC surface was defined from the depth profile of the evoked LFP. For each channel depth, we averaged the aligned LFP across repetitions of the effective stimulus and then averaged pairs of channels at the same depth from the two columns, yielding one LFP trace per depth. For each depth, we computed the peak amplitude of the visually evoked LFP. Moving from the channel with the largest amplitude to superficial channels, the surface was assigned to the first channel whose evoked LFP amplitude fell below 25% of the peak amplitude.

To locate the SGS-SO border, we computed the current-source density (CSD) as the second spatial derivative of the stimulus-aligned LFP traces and averaged those across repetitions. At the latency with maximal positive CSD amplitude, we identified the first channel below this maximum whose CSD value was negative; this depth was taken as the SGS-SO boundary.

Response amplitudes to visual stimuli

The calculation of response amplitudes in response to drifting gratings has been described previously (Schröder et al., 2020 [DOI](#)). Here, we used the same strategy for responses to static gratings and moving bars. From calcium traces, we quantified response magnitudes to drifting gratings by iteratively fitting a temporal response kernel that is 10 s long starting at stimulus onset, and a scaling factor for each presentation of a grating. First, we used a General Linear Model to fit one temporal kernel (with absolute maximum of 1) to all trials using the currently estimated scaling factors (initially set to 1). Second, we used the currently estimated temporal kernel and fitted a scaling factor for each trial, again using a General Linear Model. Both steps were repeated until the values of the estimated scaling factors stopped changing across iterations. We used the scaling factors as response strength in each trial. To estimate temporal kernels and amplitudes in response to moving bars, we introduced a third step in the iterative fitting procedure to determine the delay of the response relative to stimulus onset, which on motion directions and the location of the RF. The delays were estimated for each motion direction using the cross-correlation between the trial-averaged response and the trial-averaged prediction from the current response kernel and trial amplitudes.

To test responsiveness of a unit to a stimulus type, i.e., whether it had a stimulus response that was significantly different from baseline, we randomly shifted the response of each unit against the stimulus presentation times. A unit was determined to be *responsive* if the mean square error (MSE) of the fits to the measured responses was smaller than the 95% confidence interval of MSEs resulting from fits to the randomly shifted responses.

To estimate response amplitudes from electrophysiological recordings, we calculated firing rates from spikes between 0 and 2 s after stimulus onset and, for each trial, subtracted the firing rate between 0 and 0.5 s before stimulus onset. The unit was deemed *responsive* to a stimulus type if the grand mean response was different from zero (using a linear model and a t-test on the intercept) or if responses to different stimuli of the same type were significantly different from each other (using a linear model and an F-test on the coefficients for all stimuli).

Direction and orientation preferences and selectivity index

The direction and orientation selectivity index of a unit was determined from its mean response amplitudes, R_k , to each stimulus (drifting gratings, static gratings, moving bars). Mean amplitudes of units suppressed by gratings (responses to most stimuli <0) were inverted; remaining negative amplitudes were set to zero. Direction selectivity was assessed by scaling unit vectors pointing into the stimulus direction, α_k , by the mean amplitudes, then summing these vectors: $R = \frac{\sum_k R_k e^{i\alpha_k}}{\sum_k R_k}$. The angle and length of R are preferred direction and direction selectivity index (DSI). To determine preferred orientation and orientation selectivity, α_k was first doubled. Significance of direction and orientation selectivity, respectively, was determined by a permutation test (1,000 permutations).

Colormaps for the visualization of preferred directions and orientations are perceptually uniform (Kovesi, 2015 [DOI](#)).

Mapping of receptive fields

We presented sparse sequences of white and black squares, or white and black circles of varying sizes, to characterize the receptive fields, using the responses to white squares or circles for the ON subfield and to black squares or circles for the OFF subfield. For imaging data, we first cleaned the calcium traces by removing any slow decays at the start of experiments (likely introduced due to bleaching) and high-pass filtering the data (subtracting smoothed traces treated with a moving median with a window size of 20 s). We used linear regression to simultaneously fit two spatio-

temporal filters—one for the ON field and one for the OFF field. The time dimension of the filters spanned 0.2–0.5 s after each stimulus frame for two-photon data, and 0–0.2 s for electrophysiology data. For the ON field, only the appearance of white squares/circles was considered, for the OFF field only black squares/circles. The fits were regularized by imposing spatial smoothness on the two receptive fields (Smyth et al., 2003). We then fitted 2D Gaussians to the ON-, the OFF- and the mean between the ON- and OFF-subfield. The final spatial RF was the Gaussian that minimized the mean squared error with the mapped RF (if only one subfield was fitted by the Gaussian, the fit for the other subfield was set to zero for each pixel). The final temporal filter of the RF was determined by the least-squares scalar weights of the fitted 2D Gaussian for each time point.

A unit was considered to have a genuine receptive field if (1) the explained variance for the response to the visual noise predicted by the final spatio-temporal RF (Gaussian weighted in time) was ≥ 0.01 , and (2) the peak of the fitted Gaussian was at least 5 times the SD of the residuals between mapped RF and fitted Gaussian.

Relating RF locations to retinotopy

To relate the positions of imaged units within the FOV to their RF locations in visual space, we estimated a retinotopic transform for each recording. We modeled azimuth and elevation of the RF center as linear functions of the unit's position in the imaging plane and fitted this model under the constraint that azimuth and elevation axes in brain space were orthogonal.

We only considered units with a genuine RF (see RF mapping) and a recording had to have at least 10 such units. Units with outlier RF locations (elevation or azimuth was >5 median absolute deviations away from the median across the population) were excluded. We then fitted a linear mapping from brain coordinates (x , y) to RF azimuth az and elevation el :

$$az = a_1 + b_1x + c_1y$$

$$el = a_2 + b_2x + c_2y$$

To enforce orthogonality between the azimuth and elevation axes in brain space, we constrained the elevation coefficients (b_2 , c_2) to be orthogonal to the azimuth coefficients (b_1 , c_1):

$$b_2 = 1/\sqrt{1 + (b_1/c_1)^2}$$

$$c_2 = (-b_1/c_1)/\sqrt{1 + (b_1/c_1)^2}$$

Model parameters were obtained by weighted least squares, weighting each unit by the variance explained by its final spatio-temporal RF model, so that units with more reliable RF estimates contributed more strongly to the fit. The resulting transform defined the optimal scaling, translation, and rotation that minimized the distance between mapped RF locations and the fitted retinotopic coordinates for each recording.

Geometric model of direction and orientation preferences

To relate measured preferences to the retinal topographic organization of cardinal directions and orientations, we used geometric models derived from retinal recordings (Laniado et al., 2025 [↗](#); Sabbah et al., 2017 [↗](#)). For each unit, we computed four predicted directions and four predicted orientations at its RF location, corresponding to the four cardinal motion classes and four cardinal orientation classes defined in these studies.

Both retinal studies expressed optimal axes in retinal coordinates and then transformed them into body coordinates based on measurements of eye and head position. Axes for the left eye were provided (personal communication from Shai Sabbah) as points on the unit sphere in azimuth-elevation coordinates, where elevation is 0° at the horizon and increases towards the top of the visual field, and azimuth is 0° at the vertical meridian (nose) and increases clockwise. For motion directions, the four axes pointed to the convergence points of directional vectors on the sphere (advance, retreat, rise, fall): advance = $(180^\circ, -15^\circ)$, retreat = $(0^\circ, 11^\circ)$, rise = $(-67^\circ, -83^\circ)$, fall = $(106^\circ,$

88°). For orientations, we used four axes defining two longitudinal (“long”) and two latitudinal (“lat”) fields. Longitudinal orientations: horizontal = (187°, 90°), vertical = (98°, 26°); latitudinal orientations: horizontal = (74°, 22°), vertical = (232°, 85°).

These axes were defined under the assumption that the mouse head is pitched downward by 29° in the ambulatory posture (0° defined by bregma and lambda at the same height). Because head-fixed mice were held closer to 0° pitch, we rotated all axes about the interaural (y) axis by an angle α (0–29°) to simulate a more downward tilted head. For each mouse, we optimized α by minimizing the discrepancy between predicted and measured direction and orientation preferences. Rotation on the sphere proceeded as follows:

1. Each axis with azimuth T_{az} and elevation T_{el} was converted to Cartesian coordinates:

$$T = (\cos(T_{el}) \cos(T_{az}), \cos(T_{el}) \sin(T_{az}), \sin(T_{el}))$$
2. A rotation by angle α around the +y axis (right-hand rule) yielded:

$$U = (T_x \cos(\alpha) + T_z \sin(\alpha), T_y, -T_x \sin(\alpha) + T_z \cos(\alpha))$$
3. The result was then transformed back to azimuth-elevation coordinates:

$$T_{az} = \tan^{-1}(U_y, U_x), T_{el} = \sin^{-1}(U_z)$$

We obtained the predicted local direction or orientation at a given RF location as follows:

1. We converted the rotated axis (T_{az}, T_{el}) and the RF position (r_{az}, r_{el}) to Cartesian vectors:

$$T = (\cos(T_{el}) \cos(T_{az}), \cos(T_{el}) \sin(T_{az}), \sin(T_{el}))$$
and

$$r = (\cos(r_{el}) \cos(r_{az}), \cos(r_{el}) \sin(r_{az}), \sin(r_{el}))$$
2. For a longitudinal field, the local direction vector v at r was the component of T tangential to the sphere:

$$v = T - (T^\top r) r$$
For a latitudinal field, v was given by the cross-product:

$$v = r \times T$$
3. We then projected v onto the tangent plane at r using basis vectors aligned with increasing azimuth and elevation:

$$e_\phi = (-\sin(r_{az}), \cos(r_{az}), 0)$$

$$e_\theta = (-\sin(r_{el}) \cos(r_{az}), -\sin(r_{el}) \sin(r_{az}), \cos(r_{el}))$$

$$w = (v^\top e_\phi, v^\top e_\theta)$$
4. We computed the local direction angle x on the sphere:

$$x = \tan^{-1}(w_2/w_1)$$
5. Finally, we converted x to our convention (0° = rightward, 90° = downward) as:

$$z = 2\pi - x$$

Using these constructions, we obtained, for each unit and each model (direction and orientation), the predicted vectors corresponding to the four cardinal direction classes and four cardinal orientation classes at the unit’s RF. For each unit, we then assigned the predicted direction and orientation whose vector was closest (smallest angular difference) to the measured preferred direction and orientation. This angular difference defined the prediction error between the unit’s preference and the geometric model. To test whether the retinal geometric model explained the data better than expected by chance, we compared the median angular difference across units to a null distribution obtained by permuting RF locations 1,000 times and report the median together with the 2.5th–97.5th percentile interval of this permutation distribution.

Quantification and Statistical Analysis

All tests were two-sided, unless otherwise stated

Permutation tests were performed by generating surrogate datasets from the measured data, computing the test statistic for each surrogate dataset, and comparing the resulting null distribution to the test statistic of the measured data, i.e. whether or not the measured test statistic falls inside the 2.5–97.5 percentiles of the null distribution. The surrogate datasets were generated by randomly permuting the relevant feature/label across sample points.

Supplementary Figures

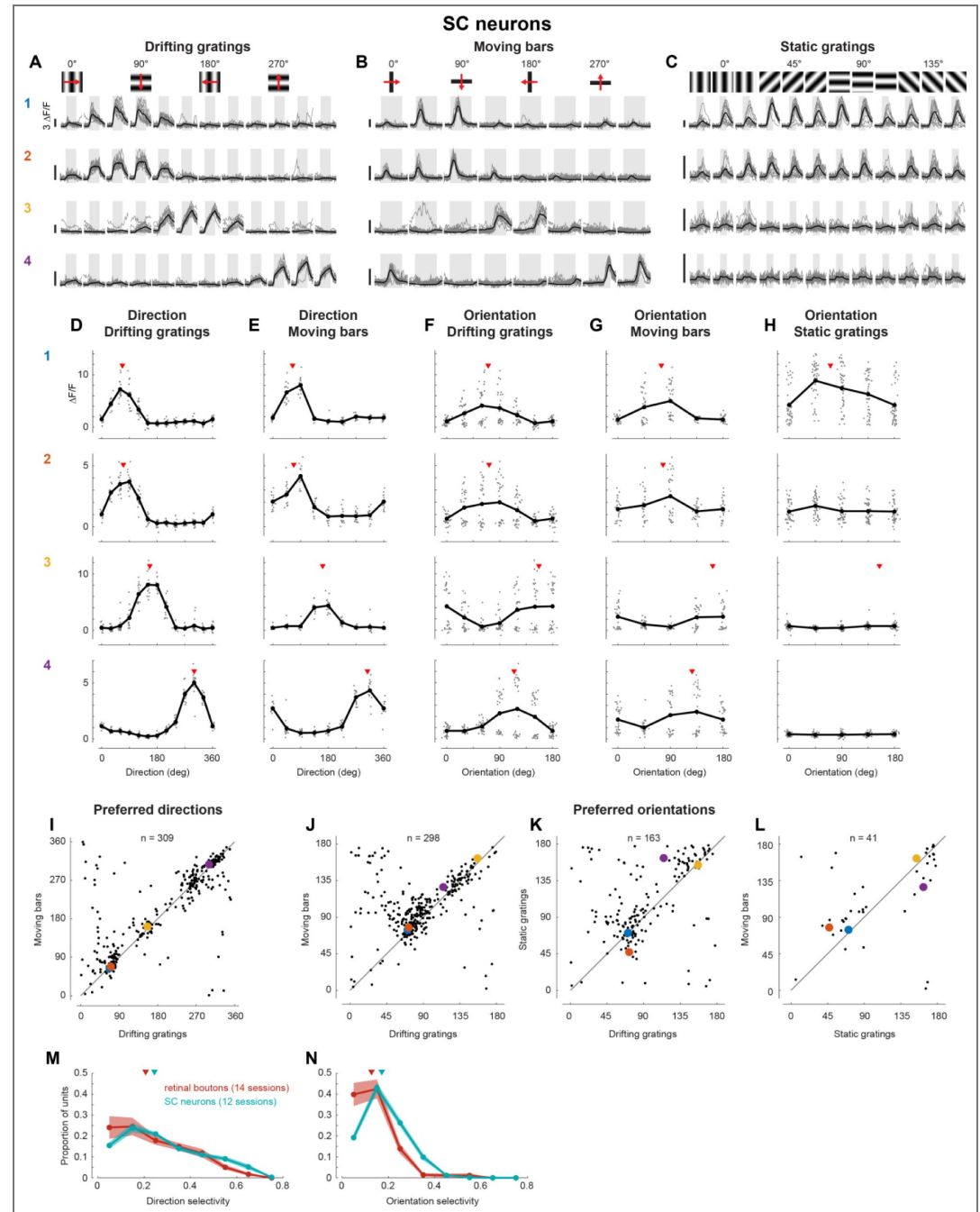


Figure S1. Direction and orientation tuning of SC neurons were consistent across stimulus types. A–C.

Single-trial (gray) and mean fitted (black) calcium traces of four example neurons in response to drifting gratings (A; 12 directions, 2 s each), moving bars (B; 8 directions, 4 s each), and static gratings (C; 4 orientations $\times$ 3 spatial phases, 1 s each). Stimulus epochs are indicated by gray shading. We recorded 2,283 SC neurons in response to moving bars and 2,314 neurons in response to static gratings, both within 6 sessions across 5 mice. As for drifting gratings, the kernel fits matched the data very well (mean $\pm$ SD explained variance: 0.33 ± 0.21 for bars, 0.30 ± 0.20 for static gratings). Across SC neurons, responsiveness dropped from 50% for drifting gratings to 31% (697/2,283) for moving bars and 41% (945/2,314) for static gratings. D–H. Direction (D, E) and orientation (F–H) tuning curves for the example neurons in A–C, showing single-trial response amplitudes (gray dots), mean amplitudes (black dots), and preferred direction or orientation (red triangles) when tuning was significant. Tuning is shown for drifting gratings (D, F), moving bars (E, G), and static gratings (H). 70% (488/697) of neurons responsive to moving

bars were selective for direction (compared to 66% for drifting gratings), while 63% (437/697) were selective for orientation (compared to 61% for drifting gratings). In contrast, only 32% (307/945) of neurons responding to static gratings were selective for orientation. I–L. Direct comparisons of preferred directions (I) and preferred orientations (J–L) obtained from different stimulus classes. Colored dots highlight the example neurons from A–C. SC neurons that were significantly tuned in response to several visual stimuli exhibited highly coherent preferences for direction and orientation with an average difference of 5.0° in preferred direction for drifting gratings and moving bars, and average differences of up to 7.0° in preferred orientation (drifting gratings vs bars: 5.3° ; drifting vs static gratings: 7.0° ; bars vs static gratings: 1.5°). M, N. Mean distribution ($\pm$ SEM) of direction (M) and orientation (N) selectivity measured with drifting gratings, across all sessions with at least 20 selective units. Triangles mark the median (across sessions) of the median selectivity index.

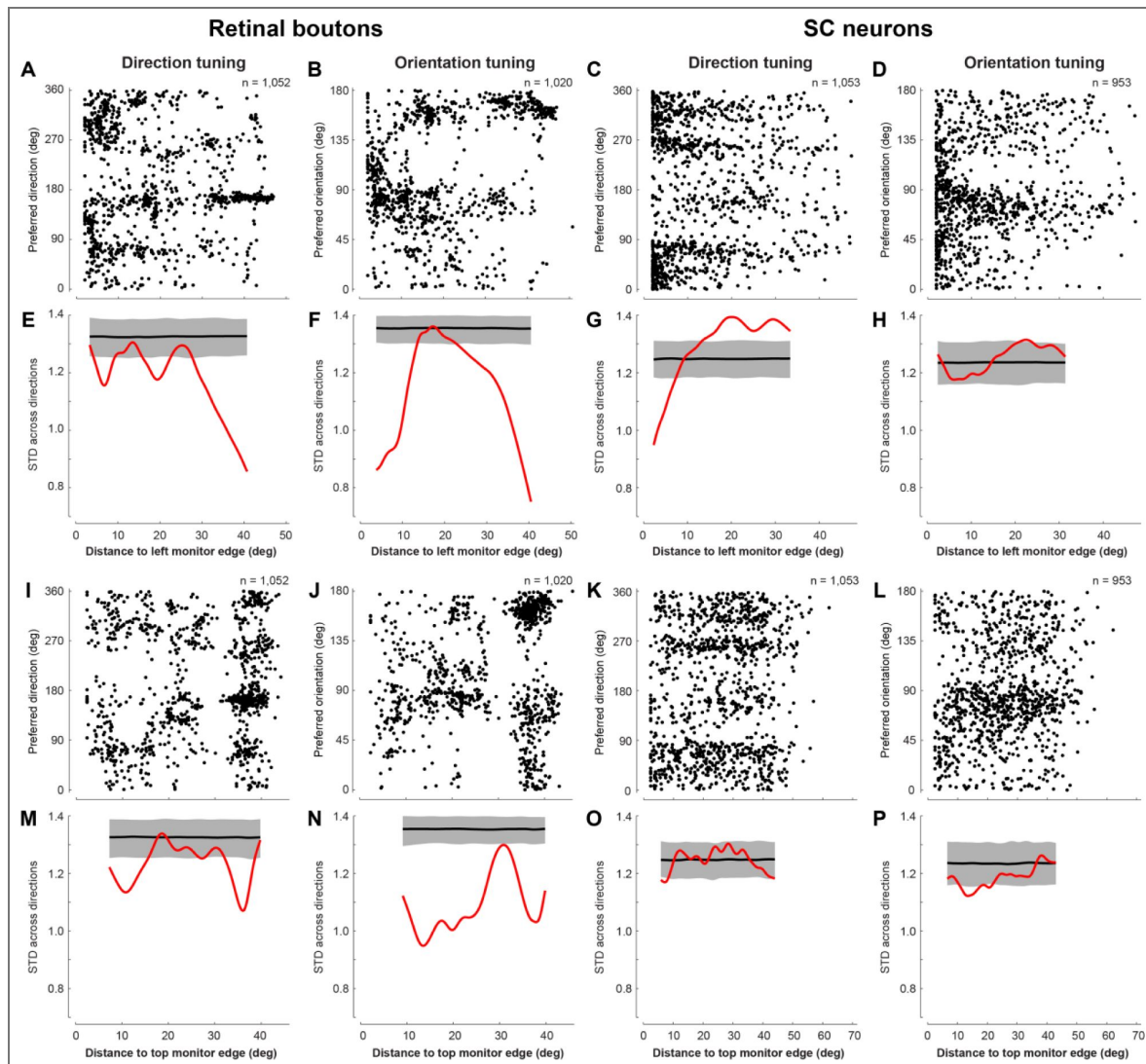


Figure S2. Proximity to monitor edges did not systematically affect measured tuning preferences.

A–D. Direction (A) and orientation (B) preferences versus distance from the RF center to the nearest vertical (left/peripheral) monitor edge, for all selectively tuned retinal boutons with mapped RFs. Corresponding data for SC neurons in C and D. E–H. Circular SD (red) of preferred directions (E) and orientations (F) for retinal boutons as a function of RF distance to nearest vertical monitor edge, compared to a null distribution obtained by permuting edge distances (black line: median; gray band: 2.5th–97.5th percentile interval). Corresponding data for SC neurons in G and H. If the monitor edge had biased tuning preferences towards a specific direction or orientation, we would expect that preferences of units closer to the monitor edge are more similar to each other than of units farther away from the edge, which was not the case in our data. I–P. As in A–H, but using distance from the RF to the nearest horizontal (top) monitor edge.

Figure S3. Topography for DS-only and OS-only units is similar to overall population.

A–D. Direction (A) and orientation (B) preferences of DS-only and OS-only retinal boutons across visual space (RFs mapped, not retinotopic). Arrows in A indicate the predicted DSGC longitude directions; colored lines in B indicate predicted OSGC longitudes and latitudes (see **Error! Reference source not found.**A–C). Numbers indicate population size in each polar histogram. Corresponding data for SC neurons in C and D. E–H. Classification of all DS-only (E) and OS-only (F) retinal boutons according to the closest predicted direction or orientation vector. Each dot shows one recording session (dot size reflects total number of boutons); black line indicates median proportion across sessions. Arrows and lines at top show example predicted vectors from a central location in A and B. Corresponding data for SC neurons in G and H. I–L. Angular differences between measured preferences and the closest predicted directions (I) or orientations (J) for DS-only and OS-only retinal boutons. Median differences (triangles) are compared to null distributions obtained by permuting RF locations (dark gray; median and 2.5–97.5 percentile intervals) or by drawing preferences from a uniform distribution (light gray; median and 2.5–97.5 percentile intervals). Direction: 9.6° original, 10.9°–12.5° permuted, 20.3°–24.4° uniform; orientation: 7.7° original, 7.9°–10.8° permuted, 12.2°–17.1° uniform. Corresponding data for SC neurons in K and L. Direction: 11.1° original, 11.7°–14.0° permuted, 19.7°–25.2° uniform; orientation: 9.1° original, 7.4°–10.2° permuted, 11.5°–17.2° uniform. Direction preferences of retinal boutons matched model predictions significantly better than those of SC neurons ($p = 0.0108$, Wilcoxon rank sum test). Orientation preferences matched model predictions to equal degrees in retinal boutons and SC neurons ($p = 0.49$, Wilcoxon rank sum test).

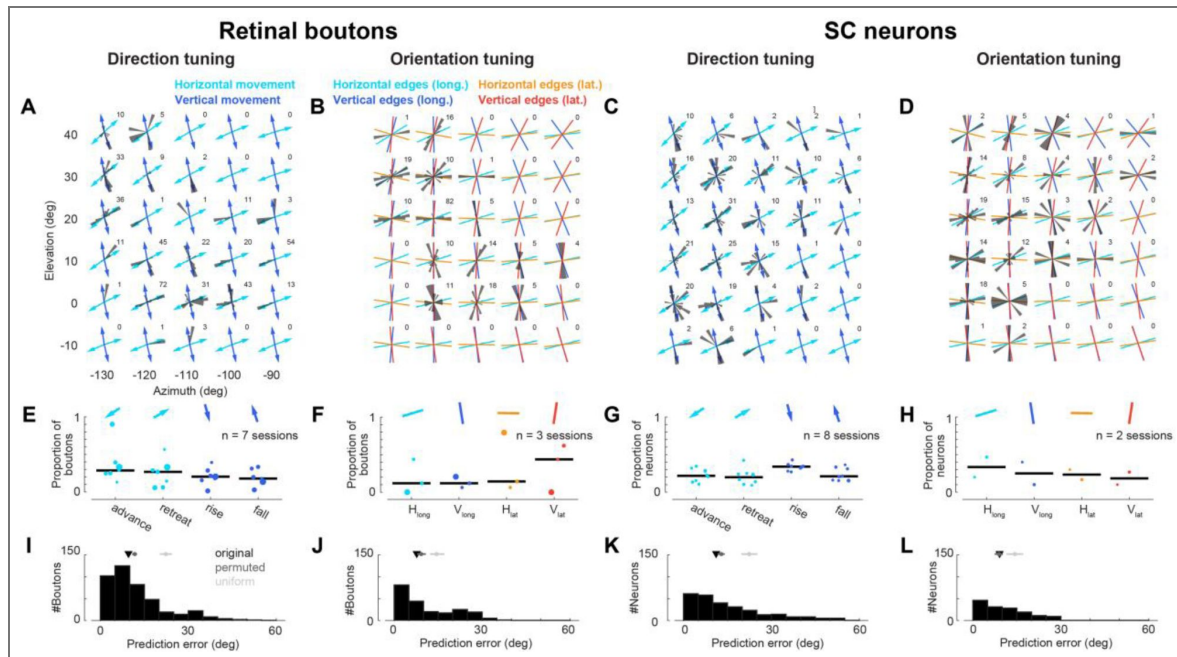
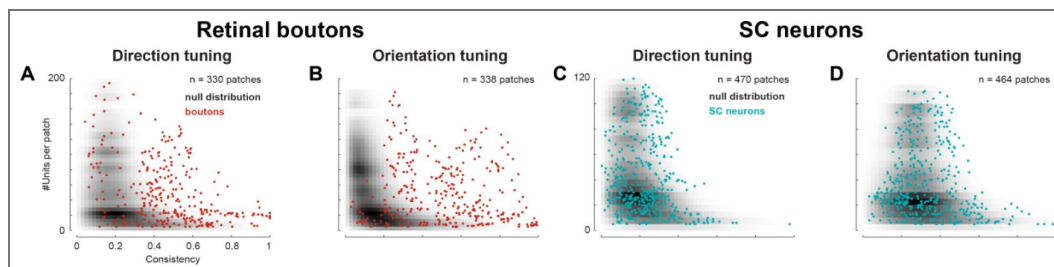


Figure S4. Local consistency is not trivially explained by the number of sampled units.

A–D. Mean consistency of direction (A) and orientation (B) preferences within each 10° patch, plotted versus the number of retinal boutons in that patch (red dots). The background grayscale indicates the density of units from a null distribution obtained by permuting RF locations across boutons. Corresponding data for SC neurons in C and D.



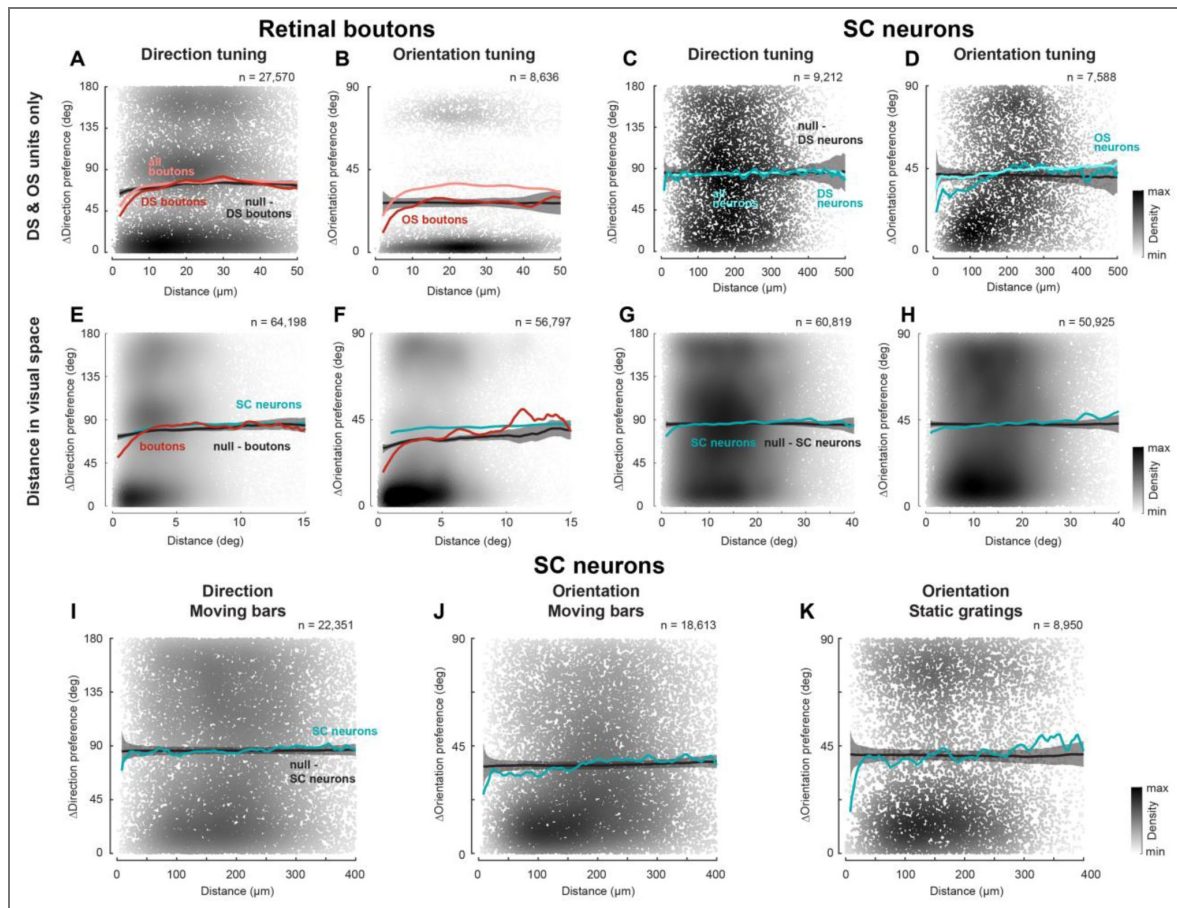


Figure S5. Weak clustering persisted within strongly selective units, with RF-based distances, and under different stimulus conditions.

A–D. Pairwise differences in direction preference for DS-only units (A) and orientation preference for OS-only units (B) versus lateral distance in SC (depth ignored) for retinal boutons. For comparison, mean pairwise differences for all tuned boutons (pink) are overlaid (same data as in Figure 5E, F). Corresponding data for SC neurons in C and D. E–H. Pairwise difference in direction (E) and orientation (F) preferences versus distance between mapped RFs for all pairs of simultaneously recorded retinal boutons. For comparison, mean pairwise differences for SC neurons (teal) are overlaid (same data as in G, H). Corresponding data for SC neurons in G and H. Only retinal boutons with RFs closer than 2.2 visual degrees had direction and orientation preferences that were more similar than expected from the null distribution. Tuning differences for SC neurons were much closer to the null distribution and were only significantly smaller for neurons with RF distances of up to 5 and 11 visual degrees for direction and orientation preferences. I. Pairwise difference in direction preferences versus lateral distance in SC (depth ignored) for all pairs of simultaneously recorded SC neurons, with preferences measured from responses to moving bars. Mean pairwise tuning differences (85°) were close to those expected from uniform distributions. J, K. Same as in I, for orientation preferences measured from moving bars (J) and static gratings (K). Mean pairwise tuning differences were close to those expected from uniform distributions (37° for moving bars, 41° for static gratings). In all panels, dots are colored by density, and mean difference as a function of distance is compared to null distribution obtained by permuting neural or RF locations (black line: median; gray band: 2.5th–97.5th percentile interval).

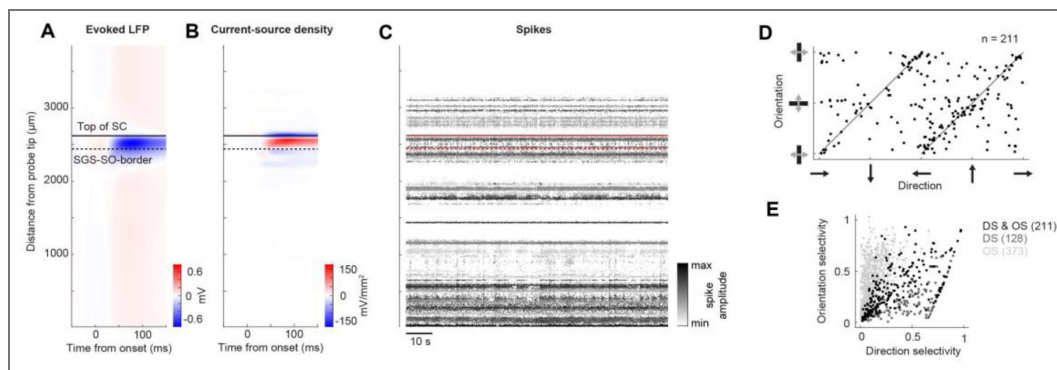


Figure S6. Determining position of SC along Neuropixels probe and tuning properties in electrophysiology data.

A, B. Evoked local field potential (LFP, A) and current-source-density (CSD, B) profiles from an example recording in response to the most effective square of the visual noise stimulus. The surface of SC is defined at 25% of the peak amplitude in the evoked LFP. The bottom of sSC (border between SGS and SO) is defined as the zero-crossing below the largest positive CSD peak. C. Spike amplitudes at spike times of all single units in the same recording as in A and B, plotted against their depths. D. Preferred direction versus orientation for SC units selective for both features. E. Direction versus orientation selectivity for all tuned SC units.

Data availability

The pre-processed data used in this study are available at <https://doi.org/10.6084/m9.figshare.30927041>; code used to analyze pre-processed data is available at https://github.com/Schroeder-Lab/he-et-al_2025_tuning-topography. The raw data are available on reasonable request.

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Additional information

Author contributions

Conceptualization and design: S.S.; data collection – two-photon imaging: S.S.; data collection and pre-processing – electrophysiology: M.F.G.F., R.G.; data analysis: S.S. and Z.H.; writing: S.S.; review and editing of manuscript: Z.H., M.F.G.F., R.G.

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Peer reviews

Reviewer #1 (Public review):

Summary:

When contemplating the role of any sensory area in the brain, an essential question is: How much of the neural code is inherited from the inputs versus constructed de novo by the local circuitry? This study tackles that important question for the case of the mouse superior colliculus (SC), a visual brain area that receives direct input from the retina. The specific aspects of the neural code are the representation of line orientation and direction of motion in the visual image. Over the past 10 years or so, there have been reports that the preferred directions and orientations of neurons vary systematically across the SC in a global map that is not present in the retina, and therefore computed locally.

Here, the authors revisit this question by expanding the range of measurements: They record from the axonal boutons of retinal ganglion cells in the input layer of the SC, from the post-synaptic neurons there, and from neurons in deeper layers of the SC. They conclude that at any given location in the SC, the signals in retinal boutons recapitulate the tuning of retinal ganglion cells, and that SC neurons follow that organization, though it is lost in the deeper layers. Notably, they find no evidence for a global map of these response properties other than what is contributed by retinal input.

Strengths:

The study combines multiple recording methods - calcium imaging and electrical recording - to capture the activity of retinal inputs to the colliculus, the tuning of neurons in the superficial layers close to the input, as well as neurons in deeper layers. Furthermore, the work connects to the recent literature on gradients of tuning properties among retinal ganglion cells. All these set the stage for testing the correspondence between retinal inputs and collicular outputs.

Weaknesses:

The methods used to identify direction-selective and orientation-selective neurons based on visual responses are overly permissive and don't match common usage in this research area. Furthermore, the measurements covered only a small fraction of the visual field, which limits their ability to distinguish between different hypotheses for the global map of visual response properties. Relatedly, the claim that retinal input patterns explain much of the layout in the superior colliculus should be made more quantitative.

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Reviewer #2 (Public review):

In this study, the authors investigate the spatial organization of direction and orientation selectivity in the mouse superior colliculus (SC) and its retinal inputs. By combining two-photon imaging of retinal boutons and SC neurons with Neuropixels recordings, they assess whether tuning preferences form structured maps or are arranged in a salt-and-pepper

fashion. They further compare SC tuning organization to previously described retinal geometric models to determine the extent to which collicular responses inherit retinal topography. The authors conclude that SC inherits a cardinally biased topographic scaffold from the retina, which progressively weakens with depth, and that strong global maps are absent.

A major strength of the study is the impressive combination of methodologies, including imaging of retinal boutons, imaging of SC neurons, and large-scale electrophysiological recordings across SC depth. The direct comparison to retinal geometric models is particularly valuable, as it situates the SC within a broader framework of retinotopic information transfer and advances our understanding of how retinal computations are transformed in downstream targets.

A limitation of the study, however, is that the imaging experiments sample only a relatively small and spatially homogeneous region of the visual field, whereas the electrophysiological recordings cover a different portion of SC. This separation makes it difficult to form a fully integrated, global picture of the spatial organization of direction and orientation selectivity across the entire collicular map.

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Reviewer #3 (Public review):

Summary:

The authors studied the organisation of orientation and direction-selective retinal ganglion cells' boutons in the mouse superior colliculus. They confirmed the results already published (Molotkov, 2023; de Malmazet, 2024; Vita, 2024; Laniado, 2025), that retinal ganglion cells' boutons in the superior colliculus conserve the retinal organisation. Thereby, orientation and direction preferences of retinal boutons at each collicular location reflect the tuning of retinal ganglion cells found at the corresponding retinal location, that is covering a matching receptive field location.

The authors also studied the organization of orientation and direction-selective neurons in the superior colliculus. They report a lack of functional organisation in the superior colliculus for neurons preferring the same stimulus orientation or direction of movement. This goes against several published reports (Ahmadlou and Heimel, 2015; Liang et al., 2023; De Malmazet et al., 2018; Feinberg and Meister, 2014; Kasai and Isa, 2021; Li et al., 2020) but echoes a study from Chen et al. (Chen, 2021). The latter authors contested the strength of the anatomical clustering of tuned alike direction-selective neurons. They found, however, that in about a quarter of their recordings, direction-selective cells with similar preferred directions did cluster anatomically in the superior colliculus.

Here, the authors of the current manuscript under review report that local clustering of tuning was weak in all neural populations and confined to very small spatial scales (10-20 μm). This is one order of magnitude smaller than previously reported clusters of around 100-300 μm wide. Therefore, the authors conclude that orientation and direction tuning in the mouse superior colliculus follows a salt and pepper organisation.

Strengths & Weaknesses:

Although the authors performed a solid analysis contesting the functional clustering of direction and orientation selective neurons, there seemed to be some elements in their data in favour of a functional clustering of neurons.

As an illustration, the authors plotted in Figure 1Q the distribution of preferred orientations from all their recorded orientation-selective cells. The curve shows a clear bias, indicating

that neurons preferring horizontal orientations were found two times more often than neurons encoding any other orientations. Moreover, the authors recorded all their neurons from a defined anatomical location of the colliculus, marked by the dotted rectangle in Figure 3A-C. Therefore, this suggests that orientation-selective cells in this particular collicular location are biased towards preferring horizontal orientations. This supports an anatomical clustering of tuned alike orientation-selective cells in the superior colliculus.

Similarly, Figure 1P shows a bias in the preferred directions of direction-selective neurons in the same recording area. Cells tended to respond more to upward and forward-moving stimuli. The bias is more modest than the one described above for preferred orientations. However, it still seems significant. For example, cells preferring upwards movements appeared to be four times more abundant than cells preferring downward movements. As a consequence, it indicates that preferred directions might not be uniformly distributed and equally represented across the superior colliculus.

These anatomical biases are also visible in the receptive field analysis of the paper. In Figure 3G, the authors plotted the distribution of preferred orientations for every 10-degree bins within the recorded field of view. Out of 26 bins containing more than one neuron, only 6 seemed to include cells not overwhelmingly preferring a single orientation. These were located towards the top right of the figure. Therefore, over almost 80% of the recorded superior colliculus, the data seem in agreement with the view that orientation-selective cells tend to prefer the same orientation at a given receptive location.

The patch analysis in Figures 4G and H also seems to show some degree of coherence in the preferred orientation and direction of neighbouring tuned collicular cells. In both Figures 4G and H, clear patches of similar preferred orientation and direction appeared to emerge. For example, in Figure 4H, there is a predominance of horizontally tuned patches. This was expected given the recording bias consisting of a majority of horizontally tuned cells. In addition, vertical and 45-degree patches are also visible, in blue and red, respectively. These patches overlap with the corresponding retinotopic locations in Figure 3G, where the histograms show that cells tend to prefer the same orientations, horizontal, vertical or 45 degrees.

It is important to note that in the previous studies on functional clustering of orientation and direction, variability in the tuning of cells within clusters was always reported (Ahmadlou and Heimel, 2015; Chen et al., 2021; De Malmazet et al., 2018; Feinberg and Meister, 2014; Kasai and Isa, 2021; Li et al., 2020). This was more marked for direction-selective cells than for orientation-selective cells. In general, cells preferring all four cardinal directions were often recorded at any given collicular location. Similarly, orientation-selective cells could be found to prefer deviant orientations compared to adjacent cells. Therefore, it is not surprising to see locally mixed tuning in collicular neurons. However, what appeared significant in these studies was the overall proportion of cells with similar tuning in patches of the superior colliculus. As described above, this also seems to be the case in the data of this manuscript.

To conclude, it seems that authors tend to overlook the sources of agreement between their data and previous reports showing functional clustering of cells in the superior colliculus. Instead, the authors tend to emphasise the dissimilarities and variability to put forward a contentious view on the organisation of orientation and direction selectivity in neurons of the superior colliculus. This, I fear, is detrimental to the field because it creates a sort of manufactured chaos that produces unnecessary confusion for readers who do not attentively read the manuscript. It would be valuable for the authors to consider rewriting the manuscript, acknowledging where their data, in fact, support some level of functional clustering.

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Author Response:

We thank the reviewers and editors for their thoughtful and constructive assessment. We are encouraged that the reviewers viewed the combination of retinal bouton imaging, collicular neuron imaging, and depth-resolved electrophysiology, together with the comparison to retinal geometric models, as a strength of the study. As the reviewers note, our findings are consistent with previous *in vitro* studies showing topographic organization of tuning in the retina and with recent work demonstrating the precision of retinotopic mapping from retina to superior colliculus (SC). In revision, we will refine our definition of tuning, sharpen our claims about the spatial organization across SC and its correspondence to retinal topography, and make clearer our aim of reconciling seemingly opposing findings in the literature. In addition, we will provide a detailed response to all other points raised by the reviewers.

A central point raised in the reviews concerns our definition of direction- and orientation-selective cells. We agree that relying only on statistical significance is not sufficient for our purposes, because the resulting classification can depend on recording duration and statistical power. In the revised manuscript, we will therefore introduce thresholding criteria for direction and orientation selectivity indices (DSI and OSI) in addition to significance-based testing. We will also make clearer that our primary question is which stimulus directions and orientations are represented at a given receptive field location, rather than the distribution of preferences among neurons classified as purely direction- or orientation-selective.

We will also revise the text to define more precisely what our data do and do not establish about the large-scale organization across SC. Our intended conclusion is not that we identify a novel global organization, which would require sampling a larger portion of visual space, but rather that the regions we sampled are not well explained by previously proposed global maps in which each visual field location is dominated by a single tuning preference and the same organization is conserved across individuals. Instead, our data are more consistent with a retinal organization of biases toward specific directions and orientations that vary systematically across visual space.

We will further clarify how we quantified the correspondence between our data and the previously established retinal model of direction and orientation tuning. In the current manuscript, we report the errors between model predictions and measured tuning preferences at the corresponding visual field locations. We then assess model performance by comparing the distribution of these errors with the errors obtained from two surrogate datasets: one in which the correspondence between visual field location and tuning preference is destroyed, and one in which the prior distribution of tuning preferences is assumed to be uniform. In the revised manuscript, we will make the interpretation of this comparison more explicit, so that the reported errors are clearly presented as the relevant effect-size measure alongside significance.

Finally, we appreciate the reviewers' concern that the manuscript may currently emphasize disagreement with previous studies too strongly. We will revise the Discussion to better acknowledge where our data support some degree of local bias or weak clustering, while clarifying that we do not find evidence for a robust, stereotyped global map that is consistent across animals. Our goal is to sharpen the manuscript so that it better reconciles seemingly divergent findings in the literature rather than setting them in opposition.

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