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Type-specific dendritic integration in mouse retinal ganglion cells

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Neural computation relies on the integration of synaptic inputs across a neuron’s dendritic arbour. However, it is far from understood how different cell types tune this process to establish cell-type specific computations. Here, using two-photon imaging of dendritic Ca2+ signals, electrical recordings of somatic voltage and biophysical modelling, we demonstrate that four morphologically distinct types of mouse retinal ganglion cells with overlapping excitatory synaptic input (transient Off alpha, transient Off mini, sustained Off, and F-mini Off) exhibit type-specific dendritic integration profiles: in contrast to the other types, dendrites of transient Off alpha cells were spatially independent, with little receptive field overlap. The temporal correlation of dendritic signals varied also extensively, with the highest and lowest correlation in transient Off mini and transient Off alpha cells, respectively. We show that differences between cell types can likely be explained by differences in backpropagation efficiency, arising from the specific combinations of dendritic morphology and ion channel densities.

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A cross the nervous system, the output signal of a neuron is determined by how it integrates the often thousands of synaptic inputs it receives across its dendritic arbour. However, still little is known about how dendritic integration is shaped by differences between neuron types, such as specific dendritic morphology and ion channel complement and density. To investigate type-specific dendritic integration and the key factors driving it, we here use the vertebrate retina, a model system with a clear input–output relationship that can be recorded in a dish. The retina decomposes the visual signal into ~40 feature-specific parallel channels (reviewed in ref. 5), relayed to the brain by a matching number of retinal ganglion cell (RGC) types7,8. RGCs receive their main excitatory drive from the bipolar cells (BCs), which pick up the photoreceptor signal in the outer retina. In addition, RGCs (and BCs) receive inhibitory input from amacrine cells (ACs) (reviewed in ref. 9), completing the canonical RGC input circuit. Different RGC types differ in morphology10–12, synaptic connectivity10,13, and expression of ion channels14,15.

To explain the emergence of diverse RGC functions, many previous studies have focused on the selective connectivity with presynaptic neurons in the inner plexiform layer (IPL) (e.g., refs. 10,16,17). Different RGC types arborize in specific layers of the IPL and, hence, receive synaptic inputs from distinct combinations of BC and AC types10. This spatiotemporally heterogeneous input provides the basis of type-specific feature extraction18. In addition, RGC dendrites may themselves perform complex computations and therefore contribute to the generation of specific output channels, e.g., through their dendritic geometry, and the complement, distribution, and density of passive and active ion channels (reviewed in refs. 11,19). So far, dendritic processing in the retina has been studied mainly in interneurons (e.g., refs. 20–22). Despite some theoretical work in this direction (reviewed in ref. 23), experimental evidence for type-specific dendritic computation and their biophysical mechanisms in RGCs remains limited and is restricted to a few specific types (i.e. direction-selective RGCs24,25; On alpha RGCs26).

Here, we exploit the unique structure of the IPL to isolate the contributions of type-specific synaptic input profiles from intrinsic cellular mechanisms to elucidate whether RGC types sampling from a similar input space use specific dendritic integration profiles to generate functionally diverse outputs. To this end, we studied the dendritic integration properties of four Off RGC types in the mouse retina that receive excitatory input from a highly overlapping set of presynaptic neurons. To record light stimulus-evoked signals across the dendritic arbor of individual RGCs, we used two-photon Ca\(^{2+}\) imaging. We found that these morphologically diverse RGC types differed strongly in their spatio-temporal dendritic integration properties. A biophysical model suggests that the differential dendritic integration in these RGC types arises from the type-specific combination of dendritic morphology and ion channel complement.

Results

Estimating local dendritic receptive fields in single RGCs. To study dendritic integration in different RGC types, we recorded Ca\(^{2+}\) signals in response to visual stimulation across the dendritic arbour of individual cells in the ex-vivo, whole-mounted mouse retina using two-photon imaging. For that, we injected individual RGCs with the fluorescent Ca\(^{2+}\) indicator dye Oregon Green BAPTA-1 (OGB-1) using sharp electrodes (Methods), resulting in completely labelled individual cells (Fig. 1a). After recording dendritic activity, the cells were 3D-constructed (Fig. 1b), allowing us to extract morphological parameters such as dendritic arbour area, branching order and asymmetry. To determine the cell’s dendritic stratification profile across the IPL relative to the ChAT bands, blood vessels labelled with Sulforhodamine 101 (SR101) were used as landmarks (Fig. 1a, b; Methods).

To map dendritic receptive fields (RFs) of RGCs (Fig. 1c, d), we used a binary dense noise stimulus (20 × 15 pixels, 30 µm per pixel) that was centred on the respective recording field. For each recording field (32 × 16 pixels @31.25 Hz), we extracted regions-of-interest (ROIs) along the dendrites using local image correlations (Supplementary Fig. 1a; Methods). Next, we registered the position and distance of each dendritic segment relative to the soma and extracted each ROI’s Ca\(^{2+}\) signal. To mitigate the effect of low signal-to-noise ratio in some dendritic recordings (Methods), we routinely applied automatic smoothness determination using a Linear-Gaussian Encoding framework27 to obtain reliable estimates of each ROI’s RF (Supplementary Fig. 1b, c).

OGB-1-mediated Ca\(^{2+}\) signals have been shown to allow detecting single action potentials and bursts28–30, as well as subthreshold events28, suggesting that the resulting Ca\(^{2+}\) signal is a useful proxy for membrane voltage. However, other factors, like Ca\(^{2+}\) buffering may have contributed to the recorded signal. To assess whether Ca\(^{2+}\) signal-derived RFs reflect membrane potential-derived RFs, we performed patch-clamp recordings to measure voltage and Ca\(^{2+}\) simultaneously while presenting the dense noise stimulus (Supplementary Fig. 2a). We found that the RF estimated from Ca\(^{2+}\) responses (in soma or proximal dendrite) were almost identical to those estimated from somatic voltage responses or the spike train (Supplementary Fig. 2b–d). In addition, we found that the gradient (the rate of change) of the recorded Ca\(^{2+}\) signals was linearly related to spike rate (Supplementary Fig. 3) in both tOff alpha and tOff mini cells. These results suggest that the light-evoked dendritic Ca\(^{2+}\) signals we measured largely reflect Ca\(^{2+}\) influx through voltage-gated channels and, hence, membrane depolarisation, consistent with previous findings31,24.

Finally, we overlaid the RF contours determined from the dendritic Ca\(^{2+}\) responses with the cell’s morphology (Fig. 1e–g). For each cell, we recorded different dendritic regions at various distances from the soma yielding between 40 and 232 ROIs per cell (Fig. 1h, i; cf. Supplementary Fig. 1c). This enabled us to systematically probe dendritic integration across an RGC’s dendritic arbour and link the properties of local dendritic RFs to overall cell morphology.

Recorded RGCs are clustered into four morphological types. To compare dendritic integration profiles across RGC types with overlapping excitatory inputs, we focussed on Off RGCs that stratify close to the Off ChAT band (Fig. 2a; Supplementary Fig. 4). We recorded n = 31 cells and clustered them into four morphological groups, using four morphological criteria: soma size, arbour asymmetry, arbour density difference, and arbour area following Bae et al.11 (Fig. 2; Methods). One group likely corresponded to transient Off alpha (tOff alpha) RGCs, as indicated by a large soma and dendritic area (for statistics, see Table 1) and their characteristic stratification profile (compare to 40w RGCs in the EyeWire database of reconstructed cells of the mouse retina, http://museum.eyewire.org). The second group likely represented the Off mini alpha transient type (tOff mini; ref. 7): Cells assigned to this group exhibited an IPL stratification very similar to tOff alpha cells, but had smaller somata and dendritic areas. The third group resembled the morphology of F\(^{\text{off}}\)32, exhibiting an IPL stratification profile peaking between the Off ChAT band and the outer IPL border and a small, highly asymmetrical dendritic arbour. Finally, the fourth
group displayed a similar IPL stratification profile as sustained Off alpha RGCs (1wt cells in ref. 11), but had smaller somata and arbour areas. These cells may correspond to the Off sustained (G7) RGCs identified by Baden et al. Here, we refer to them as sustained Off (sOff). In the following, for simplicity, we will refer to these morphological groups as RGC types.

**Dendritic integration profiles vary across RGC types.** Dendrites can process incoming synaptic inputs on a local and a global scale, resulting in rather compartmentalised and synchronised dendrites, respectively. To investigate whether the four Off RGC types differ with respect to their integration mode, we first assessed how the RF size changed as a function of dendritic distance to the soma. In tOff alpha cells, local RF area systematically decreased as a function of ROI distance from the soma (Fig. 3a–c; cf. Fig. 1g, i), suggesting that signals in distal dendrites of tOff alpha cells are more isolated and local than those in proximal dendrites. This was not the case in the three remaining RGC types, where RF size remained relatively constant across different positions of the dendritic arbour (Fig. 3a–c; for details, see Supplementary Statistical Analysis). In fact, proximal RFs were significantly larger in tOff alpha cells than in the other RGC types (Fig. 3d), which did not differ systematically in their RF size along their dendrite. Notably, the dendritic RFs of all four Off RGC types were clearly larger than those of Off BCs (Fig. 3c), suggesting that the dendritic RFs we observed largely result from

![Image of dendritic receptive fields](image-url)
spatial processing at the level of the RGC dendritic arbour. Together, among the recorded RGC types, dendritic signals in tOff alpha cells are the least spatially synchronised, suggesting that they process dendritic input more locally than the other types.

Synchronisation of dendrites can originate from strong backpropagation of somatic spikes to the dendrites (reviewed in ref. 36). This is not only expected to increase dendritic RF size but should also shift the RF’s centre closer towards the soma or, more precisely, the centre of the dendritic arbour (approximately centre of the cell’s total RF). In contrast, for a more isolated dendrite without backpropagation, the RF centre should roughly correspond to the respective ROI position. Therefore, we next analysed for the four RGC types the ROI-to-RF-centre distance (RF offset distance; Fig. 3e, f) as well as the direction of this offset for the four RGC types the ROI-to-RF-centre distance (RF offset sOOff, and difference measure. Scale bar: a 50 μm.

**Table 1: Morphological parameters describing the dendritic arborues of the clustered RGCs.**

|       | Arbour density difference [a.u.] | Area [10^3 μm^2] | Asymmetry [a.u.] | Soma size [μm^2] |
|-------|----------------------------------|------------------|-----------------|-----------------|
| tOff alpha | 17 | 53.2 ± 2.3 | 44.9 ± 6.3 | 322.5 ± 8.3 |
| tOff mini | 5 | 29.7 ± 1.9 | 18.9 ± 4.1 | 151.5 ± 14.8 |
| sOff | 4 | 29.0 ± 1.5 | 14.2 ± 3.7 | 204.0 ± 34.1 |
| F-miniOff | 5 | 11.5 ± 0.6 | 74.8 ± 6.2 | 102.7 ± 2.2 |

For parameter definitions, see Methods.

**Fig. 2 Anatomical clustering of recorded RGCs.** a. Top- and side-views of four reconstructed Off RGCs, one of each studied type, with IPL stratification profiles as mean (black) and for all recorded cells of that type (grey). Dashed lines indicate On and Off ChAT bands. b Cluster-dendrogram with the morphological parameters used in each clustering step and the resulting RGC groups: n = 17 tOff alpha, n = 5 tOff mini, n = 4 sOff, and n = 5 F-miniOff. Colours indicate cluster (RGC type), dot diameter represents soma area. Inset: Illustration of arbour density difference measure. Scale bar: a 50 μm.
next probed the temporal synchronisation of light responses across the dendritic arbour of the four RGC types. For that, we used a chirp stimulus that consisted of a light step followed by frequency and contrast modulations (Methods) and was presented as local (100 µm in diameters) and full-field (800 × 600 µm) version. Notably, F-miniOff RGCs did not show any reliable dendritic chirp responses, despite the same ROIs passing our RF quality threshold (Methods). This finding is consistent with earlier observations in this RGC type (cf. x2 cell of Extended Data Fig. 5 in ref. 7). Therefore, we focussed the following analysis on the remaining three RGC types.

We found that dendritic responses to the local chirp in tOff alpha and tOff mini RGCs were quite similar but differed from those in sOff RGCs (Fig. 5a–c). In the latter, local chirp responses were more sustained than those in the other two types (Supplementary Fig. 7a–c); this difference resonates with sOff cells stratifying slightly more distally (cf. Fig. 2a) and, hence, presumably receiving more input from sustained BC types16,37. When presented with the full-field chirp, tOff alpha and tOff mini RGC responses became somewhat more distinct (i.e. to the frequency modulation). This difference was not found in an earlier study7 but may be related to the fact that in the present study, light stimuli could be precisely centred on the recorded cell. In addition, all three RGC types often showed On-events that were much less frequent for the local chirp (Fig. 5a–c; Supplementary Fig. 7b, d). Similar On-events in Off cells have
also been observed in BC responses\textsuperscript{37}. In general, differences between full-field and local chirp responses were more pronounced in sOff RGCs (Supplementary Fig. 7e), suggesting that stimulus size had a larger effect on sOff cells compared to the other two types. This could be due to a stronger inhibitory surround or connections to BCs that are more strongly influenced by surround stimulation\textsuperscript{37}.

To analyse the temporal properties of dendritic integration in these cells, we quantified the correlation of local or full-field chirp responses between ROI pairs across the dendritic arbour (Fig. 5d; Supplementary Fig. 8). In all three RGC types, correlations between ROIs were higher for responses to local than to full-field chirps (Fig. 5d, e), possibly due to surround suppression of the centre excitatory inputs\textsuperscript{38}. The decorrelation observed for full-field chirps was especially pronounced in sOff cells (Fig. 5e). In tOff alpha and sOff RGCs, correlation decreased with dendritic and angular distance (Fig. 5d). In contrast to the other two RGCs, temporal correlation in tOff mini cells was largely independent of dendritic and angular distance (Fig. 5d). In addition, correlation was overall much higher, indicating that dendritic segments in tOff mini cells are temporally more synchronised (cf. Fig. 5c). In tOff alpha and tOff mini cells, lower or higher correlation coincided with smaller or larger RF overlap, respectively. In contrast, sOff RGCs displayed low correlation in their distal dendrites in the presence of highly overlapping RFs across the whole dendritic arbour. This is consistent with the above findings, which suggest that sOff cells may have stronger surround inhibition than the other two RGC types. Because response quality was similar for the RGC types (Supplementary Fig. 9) and differences in temporal correlation between RGC types persisted when applying a more stringent quality criterion (Supplementary Fig. 10; Methods), it is unlikely that they were due to systematic differences in recording quality (i.e. signal-to-noise-ratio).
Taken together, our data suggest that spatio-temporal integration is tuned across the RGC dendritic arbour in a highly type-specific manner (Fig. 5f, g). The studied RGC types ranged between two main dendritic integration profiles: The first profile featured strongly isolated dendrites (e.g. in tOff alpha) and may render the cell sensitive to fine visual stimulus structures within the cell’s RF. In contrast, the second profile featured strongly synchronised dendrites with highly overlapping RFs (e.g. in tOff mini RGCs) and may tune the cell towards robustly detecting a stimulus independent of its location within the RF.
Fig. 5 Temporal correlation across dendrites. a Exemplary response of a tOff alpha RGC to local (middle) and full-field chirp (right) recorded from three ROIs indicated on the reconstructed cell (left). Values next to the traces indicate linear correlation coefficient of the corresponding trace pair. b, c Like a, but for tOff mini (b) and sOff RGC (c). d Hexagon maps showing response correlations for local (left) and full-field chirp (right) as a function of angular distance and dendritic distance between ROIs for tOff alpha (n = 17\(\times\)12,770\(\times\)13,001 cells\(\times\)pairs for full-field\(\times\)pairs for local), tOff mini (n = 5\(\times\)6529\(\times\)6529), and sOff RGCs (n = 4\(\times\)2622\(\times\)2557). Colour encodes correlation. e 2D comparison maps for inter-ROI correlation of local and full-field chirp responses for each RGC type. Colour codes difference in correlation, with whitened areas indicating no significant difference. f, g Like e, but for the comparison between cell types for local chirp responses (f) and full-field chirp responses (g). For details, see Supplementary Statistical Analysis.

Fig. 6 Evaluating backpropagation using somatic current injections in tOff alpha and tOff mini cells. a Left: tOff alpha cell recorded electrically at the soma using whole-cell patch-clamp (shaded triangle represents electrode) while imaging dendritic Ca\(^{2+}\) signals. Right: Simultaneously recorded somatic voltage and dendritic Ca\(^{2+}\) signals in response to somatic 500-ms current injections (0.1 to 0.5 nA, \(\Delta I = 0.1\) nA); Ca\(^{2+}\) signals from dendritic regions indicated by the boxes (left). b Same as in a, but for a tOff mini cell. c Dendritic Ca\(^{2+}\) signal (as area under the curve) as a function of spike numbers generated in the soma (tOff alpha, n = 3\(\times\)55,80 cells\(\times\)ROIs for proximal dendrite\(\times\)ROIs for distal dendrite; tOff mini, n = 3\(\times\)55,80; linear regression and corresponding confidence interval shown as black line and grey shading, respectively). d Slope from the linear regression in c for proximal and distal dendrites and both RGC types (same colours as in a–c). Data are presented as mean with error bars indicating 95% confidence intervals. Except for the slopes estimated based on for proximal and distal dendritic recordings in tOff alpha cells, all the slopes are significantly different across conditions (F(3, 267) = 27.357, p < 0.0001). Tukey’s method is used for post hoc pairwise comparison. For details, see text and Supplementary Statistical Analysis.

Differences in backpropagation of spikes between RGC types. The results so far predict cell type-specific differences in the efficiency of backward propagation of somatic signals, particularly for tOff alpha and tOff mini cells. To test this prediction experimentally, we evaluated the backpropagation of action potentials by whole-cell patch-clamp recordings combined with dendritic Ca\(^{2+}\) imaging (Fig. 6). We injected 500-ms current steps (0.1 to 0.5 nA, \(\Delta I = 0.1\) nA) into the RGC’s soma while simultaneously recording somatic voltage and dendritic Ca\(^{2+}\) (Fig. 6a, b). We observed that dendritic Ca\(^{2+}\) signals increased much more strongly as a function of the number of evoked action potentials in tOff mini cells than in tOff alpha cells (Fig. 6c). In tOff mini cells, both proximal and distal dendritic Ca\(^{2+}\) signals strongly increased with somatic spike count (slope = 3.03\(\times\)1.40, \(r^2 = 0.44\times 0.31, p < 0.001\) for proximal/distal dendrites, n = 3 cells), while in tOff alpha cells, we observed much less pronounced increase in Ca\(^{2+}\) signals with spike count number (slope = 0.44/0.24, \(r^2 = 0.08\times 0.09, p = 0.034\times 0.006\) for proximal/distal dendrites, n = 3 cells; Fig. 6d).

Together, these data confirm that backpropagation can be detected with our dendritic Ca\(^{2+}\) imaging approach and suggest that backpropagation of action potentials is more efficient in tOff mini compared to tOff alpha cells, supporting the notion that the dendritic arbour of tOff alpha cells is more electrically isolated.

Simulation reveals mechanisms for type-specific dendritic integration. The dendritic integration properties of RGC types
may be influenced by morphological features, such as branching pattern, dendritic thickness and segment length, and the complement and distribution of ion channels\textsuperscript{5,9}. To understand which of these properties may explain the dendritic integration profiles we observed, we built a simple, morphology-inspired biophysical model and focused on the effects of the type-specific morphology and dendritic channel densities in tOff alpha and tOff mini cells (Fig. 7a).

To capture the morphological differences between the cell types, we first extracted morphological parameters from a published EM dataset\textsuperscript{11}. We found that for tOff alpha cells, dendritic radius decreased systematically with increasing branch order; this decrease was less pronounced in tOff mini cells (Fig. 7b). In addition, dendritic segment length increased with branch order for tOff alpha cells, while it remained constant for tOff mini cells. Based on these differences, we built a ball-and-stick model for each cell type (Fig. 7c). For our simulations, we provided either a proximal or distal input, with read-out positions at the dendritic tip and close to the soma (Methods). Based on the dendritic integration profiles of tOff alpha and tOff mini cells (cf. Figs. 3–6), we hypothesised that (i) in tOff alpha cells, forward propagation (from distal to proximal dendrites) should be stronger than backward propagation and (ii) that backpropagation should be strong in tOff mini cells (Fig. 7c).

To investigate the role of ion channel distribution on dendritic signal propagation, we systematically varied the dendritic density of Ca\textsuperscript{2+}-activated K\textsuperscript{+} channels (\(g_{kCa}\)) and voltage-gated K\textsuperscript{+}, Na\textsuperscript{+}, and Ca\textsuperscript{2+} channels (\(g_{k}, g_{na},\) and \(g_{ca}\)). Notably, the same combination of channel densities had quite different effects when applied to the two RGC morphologies (compare columns in Fig. 7d and Supplementary Fig. 11), highlighting how strongly the interplay between morphology and channel complement affects a cell’s dendritic signal propagation. We found that distinct, cell type-specific sets of ion channel densities were compatible with the experimentally derived hypotheses (Fig. 7d): For the tOff alpha cell model, intermediate \(g_{kCa}\) and high \(g_{na}\) and \(g_{ca}\) channel densities were required to generate stronger forward propagation compared to backward propagation (Fig. 7d, e). For the same channel densities, forward propagation in modelled tOff mini cell was so low that distal inputs were almost completely extinguished before reaching the proximal dendrite. In contrast, with higher \(g_{k}\) and lower \(g_{na}\) densities, tOff mini cells showed strong backward and substantial forward propagation, in line with our hypothesis (Fig. 7d, e).

Together, these results suggest that morphology alone does not explain the experimentally observed differences between the two cell types. Instead, our model indicates that differences in
dendritic channel densities may be responsible for the distinct dendritic integration profiles in RGCs.

Discussion

Here, we studied dendritic integration in four types of mouse Off RGC (tOff alpha, tOff mini, sOff, and F-mini\textsuperscript{[38]}) which have their dendrites in overlapping strata of the IPL and, hence, receive highly overlapping sets of synaptic input. Recordings of local, light-evoked dendritic Ca\textsuperscript{2+} signals and compartmental modeling revealed surprising differences between the cells’ spatio-temporal dendritic integration. What could these distinct integration rules be good for in terms of visual computations?

In tOff alpha RGCs\textsuperscript{11}, as the distance from the soma increased, RF area decreased and dendritic RFs became increasingly non-overlapping, with minimal offset between recording site and respective RF centre. In addition, activity on different dendritic branches was only moderately correlated. The more isolated, independent dendritic segments in tOff alpha cells may help them to detect line structures of visual stimuli and support visual computations relying on spatial resolution below the RF of the entire cell. This is reminiscent of what has been reported about On alpha cells, which possess nonlinear RFs and respond to patterns that contain local structures finer than the cell’s RF centre\textsuperscript{26}. In contrast, in tOff mini and sOff RGCs, RFs overlapped extensively and changed little in area, while their centres were systematically shifted towards the soma. In addition, the timing of responses was highly correlated across tOff mini dendrites, suggesting they may reliably detect stimuli independent of their location within the RF. For sOff RGCs, the temporal correlation between the activity of different dendritic branches decreased strongly for larger stimuli, suggesting that the cell’s computational properties change as a function of stimulus size. A possible mechanism for the dependence of temporal correlation on stimulus size—not only in the sOff cells—may be shunting inhibition provided by lateral AC circuits kicking in as stimulus size increases\textsuperscript{38,40}. F-mini\textsuperscript{Off} cells\textsuperscript{32} were similar to tOff mini and sOff RGCs with some particularities related to the high asymmetry of their dendritic arbour. Our morphologically inspired biophysical model revealed that morphological difference alone cannot explain these experimentally observed dendritic integration profiles; instead, distinct combinations of morphology, ion channel complements, and densities are required.

Dendritic integration rules have been studied extensively in the cortex (e.g. refs. 41–43). In the retina, mainly interneurons have been at the centre of interest: For example, it has been suggested that horizontal cells\textsuperscript{20} and A17 ACs\textsuperscript{22} provide locally computed feedback by conjuging signals within single varicosities. Likewise, starburst AC dendrites compute the direction of motion dendrite-wise by dividing their dendritic arbour into isolated sectors which contain 15–20 varicosities each\textsuperscript{44,45}. In RGCs, dendritic integration has been studied in direction-selective (DS) RGCs, where intrinsic properties of their dendritic arbour\textsuperscript{25,46} partially their asymmetry\textsuperscript{47}, as well as the spatial arrangement of their synaptic input (reviewed in ref. 48) contribute to the generation of DS output. RGCs with constant dendritic diameter and branch length across their dendritic arbour are thought to produce densely coupled dendritic regions. In these RGCs, their morphology could enable more efficient dendritic back-propagation and therefore lead to the synchronisation of dendritic signals\textsuperscript{53}. Indeed, we observed more independent dendritic regions in tOff alpha cells, but more spatially synchronised dendritic regions in tOff mini, sOff and F-mini\textsuperscript{Off} cells. In tOff mini and tOff alpha cells, their forward and backward propagation were differentially modulated by the same combinations of ion channel densities, confirming that dendritic morphology is a key determinant of dendritic signal propagation efficiency. However, our simulation results suggest that the dendritic integration properties of tOff alpha and tOff mini RGCs could not be explained by dendritic morphology alone but require dendritic ion channels in agreement with earlier simulation studies\textsuperscript{54}. One possible reason might be that for most RGCs, action potentials generated in the soma can back propagate to the dendritic arbour\textsuperscript{55}, which needs dendritic ion channels to enable the efficient backpropagation\textsuperscript{55,56}.

Our simulation results are based on highly simplified ball-and-stick models of RGC dendrites, as these allowed us to focus on the principles of dendritic integration. Obviously, these models come with several caveats and possibilities for future extensions: First, the
morphological parameters we used were extracted from EM data\textsuperscript{11}, where tissue can shrink due to chemical fixation, such that we may have underestimated the axial conductance based on dendrite diameter. Second, it has been reported that the branching pattern is an important variable for determining propagation efficiency of dendritic signals, mainly because of the diameter changes at branch points\textsuperscript{57}. Third, the density and complement of ion channels can vary along the dendrite\textsuperscript{51,58}, raising the possibility that spatially varying ion channel densities would allow for more refined control over dendritic computations. Finally, dendritic signalling is driven by the complex interaction of excitatory and inhibitory inputs (as already mentioned above) and the locations of the respective synapses, which will require more precise connectomic studies of the cell types and microcircuits in question. A more realistic model incorporating these aspects could allow additional insights into the mechanisms underlying the observed spatio-temporal dendritic integration rules.

**Methods**

**Animals and tissue preparation.** Mice used in this study were purchased from Jackson Laboratory and housed under a standard 12 h day/night cycle with 22 °C, 55% humidity. For all experiments, mice aged 5–8 weeks of either sex were used. We used the transgenic mouse line B6;129P2-Pcdha11(+/−)/Pcx (PJA, JAX 008069, The Jackson Laboratory, Bar Harbor, ME; refs.\textsuperscript{59} 59) cross-bred with the red florescence Cre-dependent reporter line Gt(ROSA)26Sortm9(CAG-tdTomato)Hze (Ai9tdTomato, JAX 00664, n = 3 animals), as alpha RGCs can be easily targeted due to their large soma size. Also for the electrophysiological recordings, we used wild-type mice (C57Bl/6, n = 6 animals). All animal procedures were approved by the governmental review board (Regierungspräsidium Tübingen, Baden-Württemberg, Konrad-Adenauer-Str. 20, 72072 Tübingen, Germany) and performed according to the laws governing animal experimentation issued by the German Government.

Mice were dark adapted ≥2 h before tissue preparation, then anaesthetised with isoflurane (Baxter, Hechingen Germany) and killed with cervical dislocation. The eyes were quickly enucleated in carbogenated (95% O\textsubscript{2}, 5% CO\textsubscript{2}) artificial cerebral spinal fluid (ACSF) solution containing (in mM): 125 NaCl, 2.5 KCl, 2 CaCl\textsubscript{2}, 1 MgCl\textsubscript{2}, 1.25 NaH\textsubscript{2}PO\textsubscript{4}, 26 NaHCO\textsubscript{3}, 20 glucose, and 0.5 l-glutamine (pH 7.4). After removing cornea, sclera and vitreous body, the retina was trimmed on an Anodisc (0.2 µm pore size, GE Healthcare, Pittsburgh, PA) with the ganglion cell side facing up and then transferred to the recording chamber of the microscope, where it was continuously perfused with carbogenated ACSF (at 35 °C and 4 ml min\textsuperscript{−1}). All experimental procedures were carried out under very dim red light.

**Loading of individual cells with calcium indicator.** To visualise blood vessels and avoiding them when filling individual RGCs, 5 µl of a 50 mM sulforhodamine-101 (SR101, Invitrogen/Thermo Fisher Scientific, Dreieich, Germany) stock solution was added per litre ACSF solution. Sharp electrodes for single-cell injection were pulled on a P-1000 micropipette puller (Sutter Instruments, Novato, CA) with resistances ranging between 70 and 130 MΩ. Oregon Green BAPTA-1 (OGB-1, hexapotassium salt; Life Technologies, Darmstadt, Germany; 15 mM in water), a synthetic Ca\textsuperscript{2+} indicator dye with high Ca\textsuperscript{2+} affinity (K\textsubscript{d} = 170 nM; Invitrogen) and comparatively fast kinetics\textsuperscript{29}, was loaded into individual RGCs using the single-pulse function (500 ms, −10 nA) of a MultiClamp 900A amplifier (Axon Instruments/Molecular Devices, Wokingham, UK). To allow the cells to completely fill and recover, we started recordings 1 h post injection.

**Two-photon imaging and light stimulation.** A MOp-type two-photon microscope (designed by W. Denk, MPI, Martinsried; purchased from Sutter Instruments/Product) as described previously\textsuperscript{68} was used for this study. Briefly, the system was equipped with a mode-locked Ti:Sapphire laser (MaiTai-HP DeepSee, Newport Spectra-Physics, Darmstadt, Germany), green and red fluorescence detection channels for OGB-1 (HQ 510/84, AHF, Tübingen, Germany) and SR101/dtTomato (HQ 630/60, AHF), and a water immersion objective (W Plan-Apochromat 20x/0.1 DIC M27, Zeiss, Oberkochen, Germany). For all scans, we tuned the laser to 927 nm, and used a custom-made software (ScanM, by M. Müller, MPI, Martinsried, and T.E.) running under IGOR Pro 6.3 for Windows (Wavemetrics, Portland, OR). Time-elapsed dendritic signals were recorded with 64 × 16 pixel image sequences (31.25 Hz). High-resolution morphology stacks were acquired using 512 × 512 pixel image stacks with 0.1 or 1.0 µm steps.

Light stimuli were projected through the objective lens\textsuperscript{66}. We used two alternative digital light processing (DLP) projectors: a K11 (Acer, Ahrensburg, Germany) or a LightCrafter E4500 MKII (Texas Instruments, Dallas, TX; modified by EKB Technologies Ltd., Israel). Both were equipped with light-emitting diodes (LEDs)—green (575 nm) and UV (390 nm)—that match the spectral sensitivities of mouse M- and S-opsins (for details, see refs.\textsuperscript{37}). Both LEDs were intensity-calibrated to range from 0.1 x 10\textsuperscript{5} (black background) to 20.0 x 10\textsuperscript{5} (white full field) photoisomerisations P \textsuperscript{u} s\textsuperscript{−1} cone\textsuperscript{−1}. The light stimulus was centred before every experiment, ensuring that its centre corresponded to the centre of the microscope’s scan field. For all experiments, the tissue was kept at a constant mean stimulator intensity level for ≥15 s after the laser scanning started and before light stimuli were presented.

Light stimuli were generated and presented using the Python-based software package QDspy (Table 2). Three types of light stimuli were used:

1. **Binary dense noise (20 x 15 matrix of 30 µm per pixel; each pixel displayed an independent, balanced random sequence at 5 Hz for 5 min)** for spatio-temporal receptive field (RF) mapping. The pixel size was chosen to be slightly smaller than the RF centre of single BCs (38–68 µm in diameter; ref.\textsuperscript{37}), allowing to estimate RGC dendritic RFs at single-BC resolution.

2. **Full-field (800 x 600 µm) chirp, consisting of a bright step and two sinusoidal intensity modulations, one with increasing frequency (0.5–8 Hz) and one with increasing contrast.**

3. **Local chirp:** like (2) but with a diameter of 100 µm.

**Simultaneous recordings of somatic voltage and dendritic Ca\textsuperscript{2+}.** For the simultaneous recording of somatic voltage and dendritic Ca\textsuperscript{2+} in response to light stimuli or current injections (0.1–0.5 nA, ΔV = 0.1 nA), we performed whole-cell patch-clamp recordings (electrode resistance, 7–15 MΩ). In addition to 200 µM of

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**Table 2. Software used and repositories for custom scripts and data.**

| Part | Description (link) | Company/Author | Item number (RRID) |
|------|-------------------|----------------|-------------------|
| ScanM (v2.04) | 2P imaging software running under IGOR Pro | Written by M. Müller (MPI Neurobiology, Martinsried), and T.E. | SCR_0003255 |
| pClamp (v10.6) | Electrophysiology Data Acquisition & Analysis Software | Wavemetrics, Lake Oswego, OR | IGOR Pro v6 |
| IGOR Pro (v6) | https://www.wavemetrics.com | IGOR Pro v6 | SCR_0016985 |
| Python (v3.6.7) | http://www.python.org/ | Written by T.E, supported by Tom Boissonnet (EMBL, Monterotondo) | SCR_00664 |
| R (v2.3) | The R project http://www.r-project.org/ | | |
| QDspy (v0.77) | Visual stimulation software https://github.com/eulerlab/QDspy | | |
| Scikit-learn (v0.20.0) | Software package for Python | | ref. 68 |
| Scikit-image (v0.14.2) | Software package for Python | | ref. 69 |
| Itsadug (v2.3) | Software package for R | | ref. 70 |
| Mgv (v1.8-24) | Software package for R | | ref. 71 |
| emmeans (1.4.4) | Software package for R | | |
| NEURON (v7.7.0) | Simulation environment for modelling individual neurons and networks of neurons | | ref. 66 |
| Custom scripts and data | | | |

http://retinal-functomics.net/
OGB-1, the intracellular solution contained (in mM): 120 K-gluconate, 5 NaCl, 10 KC1, 1 MgCl2, 1 EGTA, 10 HEPES, 2 Mg-ATP, and 0.5 Tris-GTP, adjusted to pH 7.2 using 1 M KOH. Before the whole-cell recordings, liquid junction potential of 15 mV were corrected with the pipette offset function of the Axopatch 200B amplifier (Molecular Devices LLC). To allow the dendrites to fill with OGB-1, cells were kept in the whole-cell mode for ~5 min before the start of the recordings. In some cases, we performed extracellular (cell-attached) recordings. Here, the cells were first injected with OGB-1 (as described above) and then targeted with ASCF-loaded electrodes. All electrophysiological data were digitised at 10 kHz using the pClamp software (Molecular Devices GmbH) and Bessel-filtered at 2 kHz. All Ca2+ signals were imaged at 32.15 Hz (64 × 16 pixel image sequences).

Reconstruction of cell morphologies. Directly after the recording, the complete dendritic morphology of the RGC was captured by acquiring a high-resolution stack. In case the cell was not bright enough to see all branches in detail, a second dye-injection was performed. Using semi-automatic neurite tracing62, we obtained cell skeletons of the recorded RGCs. If necessary, we de-warped image stack and traced cell, as described earlier7. All further analysis, such as the extraction of morphological parameters (see below), was done using custom Python scripts.

Relating recording positions to cell morphology. As the full dendritic morphology could not be imaged during Ca2+ recordings, recorded dendrites (i.e. regions of interest (ROIs), see below) were not necessarily well-aligned with the cell morphology reconstructed later. Based on the relative position of each recording field, a region 9 times larger than this recording field was cropped from the reconstructed morphology and z-projected. Next, the recording field was automatically aligned with this region using match templates from scikit-image (Table 2). The centre coordinates of all ROIs in that recording field were then calibrated to the closest dendritic branch based on their Euclidean distance. In rare cases, when automatic matching failed, the matching was done manually.

Morphological parameters and hierarchical clustering. To morphologically cluster the RGCs as described by Bae et al.15, we had to determine the relative position of the two ChAT bands, the dendritic plexus of the starburst ACs63. For this, the blood vessel plexus in GCL and INL served as landmarks. With their positions defined as 0 and 1, the relative IPL depth of the On and Off ChAT bands is 0.48 and 0.77, respectively, as shown earlier15,25. The following parameters were extracted for each cell:

- Dendritic RF overlap index: An RF overlap index (Oi) was calculated as follows:

\[
O_i = \frac{A_{\text{overlap}}}{A_{\text{convex hull}}}
\]

where Aoverlap corresponding the area of a region with closed contour, and Aconvex hull the 2D morphology’s convex hull. Only data with a good RF (a single contour with li < 0.1, Aoverlap > 1.8 × 1000 mm², at a contour threshold of 0.60; see Supplementary Fig. 1) were used for further analyses. The cut-off on irregularity (li) was set to ensure the RF was approximately round, while the cut-off on Aoverlap was set to a value much smaller than the RF of a RGC, that still ensured that small, noise-generated “bumps” were excluded (cf. Supplementary Fig. 1b).

- RF Offset distance and angle: RF offset distance was calculated as the linear distance between ROI centre and the geometrical centre of its RF contour. RF Offset angle was calculated as the angle between lines from ROI centre to geometrical centre of its RF contour and ROI centre to dendritic arbour centre.

- Dendritic RF overlap index: An RF overlap index (Oi) was calculated as follows:

\[
O_i = \frac{\text{Var}(C_i^2)}{\text{Var}(C_i)}
\]

where C is the T-by-R response matrix (time samples by stimulus repetitions) and (.) and Var[.] denote the mean and variance across the indicated dimension, respectively. If all trials were identical, such that the mean response is a perfect representation of the response, Qi = 1. If all trials were completely random with fixed variance, such that the mean response is not informative about the individual trials, Qi ∝ 1/R.

- Signal correlation: To quantify temporal signal correlation, we cross-correlated the mean Ca2+ responses. We noticed that some ROIs with good spatial RFs (see above) displayed low signal-to-noise chirp responses. Hence, we repeated the analysis for Qi > 0.4 or 0.5 with comparable result (Supplementary Fig. 10).

Further temporal analysis: Using the responses to the step part of the chirp stimulus, we calculated a transience index (Ti, for local chirp) and a polarity index (Qi, for both local and full-field stimuli) as marginal IPL (cf. Fig.2b, inset). The marginal-central arbour density difference, we defined as 0 and 1, the relative IPL depth of the On and Off ChAT bands

\[
\rho = \frac{A_{\text{On}}}{A_{\text{Off}}}
\]

where AOn and AOff are the RF areas of the ROI pair, AOn is the overlap area between AOn and AOff, and AOn[. AOff[.] corresponds to the smaller area (AOn or AOff). To determine the quality of spatial RFs, contours were drawn on up-sampled and normalised RF maps with different thresholds (0.60, 0.65, and 0.70). The quality was then determined from the number of regions with closed contours, their sizes and their degree of irregularity. The irregularity index was defined as

\[
I_i = 1 - \frac{A_{\text{overlap}}}{A_{\text{convex hull}}}
\]

where Aoverlap corresponding the area of a region with closed contour, and Aconvex hull the 2D morphology’s convex hull. Only data with a good RF (a single contour with li < 0.1, Aoverlap > 1.8 × 1000 mm², at a contour threshold of 0.60; see Supplementary Fig. 1) were used for further analyses. The cut-off on irregularity (li) was set to ensure the RF was approximately round, while the cut-off on Aoverlap was set to a value much smaller than the RF of a RGC, that still ensured that small, noise-generated “bumps” were excluded (cf. Supplementary Fig. 1b).

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\]

where C is the T-by-R response matrix (time samples by stimulus repetitions) and (.) and Var[.] denote the mean and variance across the indicated dimension, respectively. If all trials were identical, such that the mean response is a perfect representative of the response, Qi = 1. If all trials were completely random with fixed variance, such that the mean response is not informative about the individual trials, Qi ∝ 1/R.

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\[
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where AOn and AOff are the RF areas of the ROI pair, AOn is the overlap area between AOn and AOff, and AOn[. AOff[.] corresponds to the smaller area (AOn or AOff). To determine the quality of spatial RFs, contours were drawn on up-sampled and normalised RF maps with different thresholds (0.60, 0.65, and 0.70). The quality was then determined from the number of regions with closed contours, their sizes and their degree of irregularity. The irregularity index was defined as

\[
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\]

where Aoverlap corresponding the area of a region with closed contour, and Aconvex hull the 2D morphology’s convex hull. Only data with a good RF (a single contour with li < 0.1, Aoverlap > 1.8 × 1000 mm², at a contour threshold of 0.60; see Supplementary Fig. 1) were used for further analyses. The cut-off on irregularity (li) was set to ensure the RF was approximately round, while the cut-off on Aoverlap was set to a value much smaller than the RF of a RGC, that still ensured that small, noise-generated “bumps” were excluded (cf. Supplementary Fig. 1b).

- RF Offset distance and angle: RF offset distance was calculated as the linear distance between ROI centre and the geometrical centre of its RF contour. RF Offset angle was calculated as the angle between lines from ROI centre to geometrical centre of its RF contour and ROI centre to dendritic arbour centre.

- Dendritic RF overlap index: An RF overlap index (Oi) was calculated as follows:

\[
O_i = \frac{\text{Var}(C_i^2)}{\text{Var}(C_i)}
\]

where C is the T-by-R response matrix (time samples by stimulus repetitions) and (.) and Var[.] denote the mean and variance across the indicated dimension, respectively. If all trials were identical, such that the mean response is a perfect representative of the response, Qi = 1. If all trials were completely random with fixed variance, such that the mean response is not informative about the individual trials, Qi ∝ 1/R.

- Signal correlation: To quantify temporal signal correlation, we cross-correlated the mean Ca2+ responses. We noticed that some ROIs with good spatial RFs (see above) displayed low signal-to-noise chirp responses. Hence, we repeated the analysis for Qi > 0.4 or 0.5 with comparable result (Supplementary Fig. 10).

Further temporal analysis: Using the responses to the step part of the chirp stimulus, we calculated a transience index (Ti, for local chirp) and a polarity index (Qi, for both local and full-field chirps). Here, only ROIs with Qi > 0.4 were used for the analysis. Before the computation of these indices, the mean traces were binomially smoothed (with 3000 repetitions). Then, 2 s.d. of the baseline (2500 samples of the smoothed trace before light stimulus onset) were used to determine the time of response onset (Tstim) and offset (Tstim). Ti was calculated as

\[
T_i = T_{\text{stim}} - T_{\text{stim}}^0
\]

where Tstim is the stimulus duration.
For POi, data points before and after the response (see above) were set to zero, before calculating:

\[ POi = \frac{1}{b} \left( \sum_{t=0}^{b} r(t + t_{on}) - \sum_{t=0}^{b} r(t + t_{off}) \right) \]

Where \( b = 3 \times t_{on} \) and \( t_{on} \) and \( t_{off} \) are the time points of light stimulus onset and offset, \( r(t) \) is the mean response at time \( t \). For ROIs responding only to the light-onset, \( POi = 1 \), whereas for ROIs only responding during the light-offset, \( POi = -1 \).

Backpropagation: For each ROI, \( \text{Ca}^{2+} \) traces were normalised by the s.d. of the baseline (first 5 s of the raw trace). To compensate for changes in baseline, the mean of the 5-s trace before the onset of the current step was then subtracted from the current-evoked \( \text{Ca}^{2+} \) response. The dendritic \( \text{Ca}^{2+} \) response (area under the curve) of the peak action potential count was determined within a time window of \(-1.6 \) s, starting from the onset of the 500-ms current step.

### Statistical analysis

We used Generalised Additive Models (GAMs) to analyse the relationships of RF size vs. dendritic distance; RF offset vs. dendritic distance; RF overlap vs. dendritic distance and dendritic angle; temporal correlation vs. dendritic distance and dendritic angle (for details, see Supplementary Statistical Analysis). GAMs extend the generalised linear model by allowing the linear predictors to depend on arbitrary smooth functions of the underlying variables \( ^{65} \):

\[ g(\mu) = \beta_0 + f_1(x_1) + \cdots + f_n(x_n) \]

Here, \( x_i \) are the predictor variables, \( g \) is a link function, and the \( f_i \) are smooth functions of the predictor variables. These smooth functions can also depend on more than one predictor variable.

To implement GAMs and perform statistical testing, we employed the mgcv package for R (Table 2). Here, for smooth terms we used penalised regression splines. We modelled the dependence of our variable of interest as a single smooth term per cell type for univariate dependencies and a tensor product smooth for bivariate dependencies. The dimension of the basis was set high enough such that the estimated degrees of freedom stayed sufficiently below the possible maximum. Smoothing parameters were selected via the default methods of the package. All models also included a random effect term for cell identity. Typically, we used models from the Gaussian family; for the dependence of RF overlap on dendritic distance and dendritic angle, we instead used a scaled \( t \)-distribution, as this improved BIC (Bayesian information criterion) and diagnostic plots (see also Supplementary Statistical Analysis).

Statistical significance for differences in the obtained smooths between cell types were performed using \texttt{plot.diff} or \texttt{plot_difff} of the \texttt{itsadug} package for R (Table 2). 95% confidence intervals were calculated using the simultaneous confidence intervals (CI) option, excluding the random effect of cell identity.

To analyse the relationship between \( \text{Ca}^{2+} \) signals and somatic action potential count evoked by current injections, we used linear regression. Model fitting was performed using R’s build-in function \texttt{lm} and the \texttt{emmeans} package for R. Statistical significance for interaction of dendritic \( \text{Ca}^{2+} \) signals and somatic action potentials and pairwise comparison of slopes were performed using R’s build-in function \texttt{anova}, \texttt{pairs}, and \texttt{lstrends} (\texttt{emmeans}) (see also Supplementary Statistical Analysis).

### Biophysical model

To explore the mechanisms underlying dendritic integration in different RGC types, we built a multi-compartmental 1D model. To get precise measurements of dendrite thickness and segment length for TOff alpha and TOff mini cells, we extracted these information from published morphologies of 40w and 4i RGCs (cf. Fig. 7b), respectively, reconstructed from EM data (http://museum.eyswire.org). Then we mapped the medium values of these parameters to the respective branch order of the model (cf. Fig. 7c). The model was implemented in the NEURON simulation environment\(^ {37} \). Here, each dendritic portion (between two branch points) was represented as a section in the simulator, which was further divided into multiple segments (compartments) with a maximal length of 7 \( \mu \)m.

The 1D model can be characterised by the cable equation,

\[ \frac{d^2V}{dx^2} = C_m \frac{dV}{dt} + I_{in} - I_{sim} \]

where \( V \) is the voltage across the cell membrane, \( x \) is the distance along the cable, \( d \) is the dendritic diameter, \( r_c \) is the intracellular resistivity, and \( C_m \) is the specific membrane capacitance. \( I_{in} \) represents the sum of four voltage-gated cation currents (sodium, \( I_{Na} \); calcium, \( I_{Ca} \); delayed rectifier potassium, \( I_{K} \); A-type potassium, \( I_{K, A} \)), one calcium-activated potassium current (\( I_{Ca, A} \)), and one leak current (\( I_{Lack} \)). The current dynamics are described following Fohlmeister and Müller\(^ {67} \) as:

\[ I_{in} = I_{Na} + I_{Ca} + I_{K} + I_{K, A} + I_{Ca, A} + I_{Lack} \]

\[ \begin{align*}
  I_{Na} &= g_{Na} m^3 h(V - V_{Na}) + g_{Na} s^4(V - V_{Na}) \\
  I_{Ca} &= g_{Ca} c^{a} (V - V_{Ca}) \\
  I_{K} &= g_{K} f(V - V_{K}) \\
  I_{K, A} &= g_{K, A} a^{h} (V - V_{K}) \\
  I_{Ca, A} &= g_{Ca, A} (V - V_{Ca}) \\
  I_{Lack} &= g_{Lack} (V - V_{Lack})
\end{align*} \]

The intracellular stimulation current \( I_{in} \) in Eq. 8 was the product of a 5000 \( m \times 200 \mu \)m 1D Gaussian noise stimulus and a BCS spatial RF with a Gaussian shape (with the width set by \( \sigma = 6 \) and the centre depends on the current injection location). The stimulation current was injected either at the proximal (at 25 \( \mu \)m from soma) or distal dendrite (at 85% of the total dendrite length from soma), then filtered by a soft rectification function. The magnitude of \( I_{in} \) was scaled between 0 and 15 nA with a mean of 6.17 nA and an s.d. of 1.88 nA to ensure that the input stimulus would elicit spikes in the soma for all parameter combinations.

Model parameters (Table 3) and channel conductances (Table 4) were taken from Fohlmeister and Miller (ref. \( {67} \)). To identify parameters that explain the experimental data, we grid-searched combinations of ion channel densities by multiplying the reference parameters with different scaling factors (for \( g_{Na} \) and \( g_{Ca} \), the scaling factors ranged from 0 to 2, and were incremented by 0.25 each step). For \( g_{K} \), we used \( 0.1, 1, 2 \), for \( g_{Ca, A} \), \( 10, 100, 1000, 5000, 90000 \).

Voltage changes in the dendrites were read-out at the centre of each dendritic section and used to estimate the local dendritic RF with a maximum likelihood method, then smoothed by a Savitzky–Golay filter (window length of 31; 3rd degree polynomial). The peak amplitudes of dendritic RFs were measured and normalised to the RF at the position closest to the current input.

### Reporting summary

Further information on research design is available in the Nature Research Reporting Summary linked to this article.

### Data availability

All relevant data are available at https://doi.org/10.5281/zenodo.3708064.

### Code availability

All custom scripts in Python, R and Igor for data processing, statistical analysis, and plotting are available at https://github.com/berenslab/rgc_dendrites. Code for receptive field estimation is available at https://github.com/berenslab/RFest.

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### Table 3 Model parameters.

| Parameters          | Values          |
|---------------------|-----------------|
| Temperature         | \( T = 32 \degree C \) |
| Intracellular axial resistivity | \( R_c = 110 \, \Omega \) cm |
| Specific membrane resistance | \( R_m = 15,000 \, \Omega \) cm² |
| Specific membrane capacitance | \( C_m = 1 \, \mu \)F cm⁻² |
| Potassium reversal potential | \( V_{k} = -75 \, mV \) |
| Sodium reversal potential | \( V_{n} = 35 \, mV \) |

### Table 4 Reference distribution of ion channels in cell compartments.

| Channel type | Conductance in soma (S cm⁻²) | Conductance in dendrites (S cm⁻²) |
|--------------|------------------------------|----------------------------------|
| \( g_{Na} \)  | 0.08                         | 0.025                            |
| \( g_{Ca} \)  | 0.0015                       | 0.0002                           |
| \( g_{K} \)   | 0.018                        | 0.012                            |
| \( g_{Ca, A} \)| 0.054                        | 0.036                            |
| \( g_{Ca, C} \)| 0.0000065                    | 0.0000001                        |
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Author contributions
The study was conceived and designed by T.B, P.B., T.E., and K.F.; Y.R. carried out the two-photon imaging with input from K.F. and T.E.; Y.R. performed the electrical recordings with input from T.S. and T.E.; Z.H. performed the cell clustering, RF estimation, and modelling with inputs from P.B. and T.E.; P.B. and Z.H. performed the statistical analysis with input from H.B.; Y.R. wrote the first draft of the manuscript; Y.R., Z.H., T.B., P.B., K.F., and T.E. edited the manuscript.

Competing interests
The authors declare no competing interests.

Additional information
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