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Spectral inference reveals principal cone-integration rules of the zebrafish inner retina

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SUMMARY

Retinal bipolar cells integrate cone-signals at dendritic and axonal sites. The axonal route, involving amacrine cells, remains largely uncharted. However, because cone-types differ in their spectral sensitivities, insights into bipolar cells’ cone-integration might be gained based on their spectral tunings. We therefore recorded in vivo responses of bipolar cell presynaptic terminals in larval zebrafish to widefield but spectrally resolved flashes of light and mapped the results onto spectral responses of the four cones. This “spectral-circuit-mapping” allowed explaining ~95% of the spectral and temporal variance of bipolar cell responses in a simple linear model, thereby revealing several notable integration rules of the inner retina. Bipolar cells were dominated by red-cone inputs, often alongside equal sign inputs from blue- and green-cones. In contrast, UV-cone inputs were uncorrelated with those of the remaining cones. This led to a new axis of spectral opponency where red-/green-/blue-cone “Off” circuits connect to “natively-On” UV-cone circuits in the outermost fraction of the inner plexiform layer – much as how key colour opponent circuits are established in mammals. Beyond this, and despite substantial temporal diversity that
was not present in the cones, bipolar cell spectral tunings were surprisingly simple. They either approximately resembled both opponent and non-opponent spectral motifs already present in the cones or exhibited a stereotyped non-opponent broadband response. In this way, bipolar cells not only preserved the efficient spectral representations in the cones, but also diversified them to set up a total of six dominant spectral motifs which included three axes of spectral opponency.

KEYWORDS. Retina, zebrafish, bipolar cell, cone-photoreceptor, amacrine cell, colour vision, spectral processing

INTRODUCTION

For colour vision, retinal circuits combine and contrast the signals from spectrally distinct types of photoreceptors. For this, our own trichromatic vision uses spectral signals along two main opponent axes: “blue-yellow” and “green-red”. Of these, blue-yellow comparisons are based on ancestral cone-type selective retinal circuits that differentially contact SWS1- (“blue”) and LWS-cones (“green/red”, aka. “yellow”), while reliably contrasting “green-red” is thought to require the central brain. This is because primate “green”- and “red-cones” emerged from a relatively recent LWS gene duplication that enabled new green sensitivity in some LWS-cones, however without providing a known means for postsynaptic retinal circuits to distinguish between “green” and “red” LWS-cone variants. Accordingly, in our own eyes, one axis of spectral opponency arises in the retina, and a second is probably decoded only in the brain.
In contrast, most non-mammalian vertebrate lineages, including fish, amphibians, reptiles, and birds, retain the full complement of ancestral cone-types based on four opsin-gene families: SWS1 (UV-cones), SWS2 (blue-cones), RH2 (green-cones), LWS (red-cones). Each of these four ancestral cones provide type-specific extracellular matrix proteins that developmental programmes use to build cone-type selective circuits in the outer retina (e.g. zebrafish, chicken). Accordingly, in these non-mammalian lineages, the expectation is that up to tetrachromatic colour vision should be possible based on stereotyped cone-opponent ancestral circuits that are specified during development, without a necessity for building additional spectral opponencies in the brain. In agreement, physiological recordings from retinal neurons in cone-tetrachromatic species including turtles and diverse species of fish consistently revealed a rich complement of complex spectral signals, including diverse spectral opponencies.

However, what the dominant opponencies are, and how they are built at the circuit level remains incompletely understood in any cone-tetrachromat vertebrate. This is in part because already horizontal cells in the outer retina functionally interconnect and potentially retune cone-types, thus limiting the possibility of making inferences about spectral processing based on recordings from downstream neurons. To address this, we recently measured the in-vivo spectral tuning of the synaptic outputs from the four cone-types in larval zebrafish using spatially widefield but spectrally narrow flashes of light. This revealed that red-cones are non-opponent, green- and blue-cones are strongly opponent with distinct zero crossings (~523 and ~483 nm, respectively), and UV-cones are weakly opponent with a zero crossing at ~450 nm. Accordingly, in larval zebrafish already the cone-output provides up to three axes of spectral opponency. However, the opponent axis
provided by UV-cones was weak, which left its role in zebrafish colour vision unclear.
Moreover, in view of expected extensive mixing of cone-signals in downstream
circuits\textsuperscript{12,28}, if and how the cones’ spectral axes are propagated downstream remains
unknown.

Accordingly, we asked how downstream retinal circuits make use of the spectrally
complex cone signals to either consolidate or to retune their spectral axes for
transmission to the brain. For this, we used two-photon (2P) imaging to measure
spatially widefield but spectrally highly resolved tuning functions at the level of retinal
bipolar cell (BCs) presynaptic terminals in the inner retina. This strategy was
previously used to establish the spectral tunings of the cones\textsuperscript{27,29}, thus facilitating
direct comparison.

We find that all three spectral axes already set-up by the cones are conserved at the
level of BC presynaptic terminals, and no new axes are created. However, the “UV-
red” axis was notably boosted and diversified into numerous variants of either
polarity via new opponent circuits that derive from red-/green-/blue-Off-circuits
connecting to UV-On-circuits. The remaining non-opponent BCs were either broadly
tuned, likely built by pooling signals from all four cone types, or essentially
resembled the tunings of red- and/or UV-cones in isolation. Beyond spectral tuning,
bipolar cells showed a rich complement of temporal features that were absent in
cones, which were notably intermixed with spectral information.

Taken together, larval zebrafish BC-circuits for colour vision therefore directly built
upon the existing cone-tunings rather than set up fundamentally new opponencies,
while at the same time adding substantial temporal complexity to the retinal code.
RESULTS

A complex interplay of spectral and temporal signals amongst BCs. To establish *in vivo* spectral tuning functions at the level of individual presynaptic terminals of bipolar cells (BCs) in the inner retina, we imaged light-evoked calcium responses from 6-7 days post fertilisation (*dpf*) RibeyeA:SyjGCaMP7b zebrafish under two-photon (2P) using established protocols\textsuperscript{19,30,31} (Methods). To record from 100s of individual BC terminals in parallel, we used a non-telecentric triplane imaging approach\textsuperscript{32} (Methods). For light-stimulation, we used the same system and protocol previously employed to determine cone-tunings\textsuperscript{27} (Figure 1A,B). In brief, light from 13 spectrally distinct LEDs was collected by a collimator after reflecting off a diffraction grating which served to narrow individual LED spectra reaching the eye\textsuperscript{33}. From here, stimuli were presented to the fish as widefield but spectrally narrow flashes of light (1.5 s On, 1.5 s Off, starting from “red” and sweeping towards UV; Methods). One example recording from BC terminals is illustrated in Figure 1C-E alongside averaged cone-responses to the same stimulus (Figure 1B) taken from Ref\textsuperscript{27}. In short, each recording plane was automatically processed to detect the boundaries of the inner plexiform layer (IPL, Figure 1D, left) and to place regions of interest (ROIs) based on pixel-wise response coherence over consecutive repeats (Figure 1D, right, Methods). From here, fluorescence traces from each ROI were extracted, detrended, z-scored, and averaged over typically 7-8 stimulus repetitions (Figure 1D,E). This revealed a great diversity in both the spectral and the temporal composition of responses amongst BCs. For example, some ROIs were entirely non-opponent but differed in their spectral tuning and in the degree to which they “overshot” the baseline between stimulus presentations (Figure 1F, compare ROIs labelled BC1 and BC2). Other ROIs such as the one labelled BC3 were spectrally opponent, here
exhibiting Off-signals to mid-wavelength stimulation but On-signals to UV-stimulation. Finally, some ROIs including the one labelled BC4 exhibited different temporal responses to long- and short-wavelength stimulation.

Because stimuli were always presented in spectral sequence, which might cause systematic adaptation, we also performed a small number of control experiments with a reduced stimulus set, where we directly compared responses of the same ROIs to ordered and to pseudorandomised stimulus sequences (Figure S1A-G). This showed that both approaches gave very similar responses, suggesting that spectral adaptation was not a major feature in our recordings. We recorded responses from a total of \( n = 72 \) triplane scans in \( n = 7 \) fish, across four major regions of the eye: Acute Zone (AZ), Dorsal (D), Nasal (N), and Ventral (V). From here, \( n = 6,125 \) ROIs (\( n_{AZ,D,N,V} = 2,535, 1,172, 1,889, 529 \), respectively) that passed a minimum response quality criterion (Methods) were kept for further analysis. Next, we clustered BC responses using a mixture of Gaussian model as described previously \(^{19,22,34,35} \) (Methods). This yielded 29 functional BC-clusters (Figure 2A,B), here arranged by their mean stratification position in the IPL (Figure 2C). If and how this relatively large number of functional BC-clusters maps onto veritable BC ‘types’\(^ {28} \) remains unknown. For comparison, previous studies described 25 functional\(^ {19} \) and 21 anatomical\(^ {12} \) BCs, however a deeper census of zebrafish BC-types, for example based on additional data from connectomics\(^ {36} \) and/or transcriptomics\(^ {37} \) remains outstanding.

Consistent with previous work that was based on a different stimulus with lower spectral resolution\(^ {19} \), zebrafish BC-clusters were highly diverse, and many exhibited a regional bias to one or multiple parts of the eye (Figure 2D). For example, several UV-dominated clusters showed a clear regional bias to the acute zone (e.g. C\(^ {21,25} \))
and/or the ventral retina (e.g. C_6,27), while many broadband clusters were distributed
approximately homogeneously across the eye except ventrally (e.g. C_2,5). However,
with our current focus on BC-spectral tunings, we did not further analyse this eye-
wide regionalisation.

Overall, BC-clusters differed strongly in their wavelength selectivity. For example,
clusters C_1 and C_2 both hyperpolarised in response to all tested wavelengths, but C_2
was tuned broadly while C_1 exhibited a notable dip in response amplitudes at
intermediate wavelengths. Other clusters exhibited clear spectral opponency. For
example, clusters C_26-29 all switched from Off-responses to long wavelength
stimulation to On-responses at shorter wavelengths. A single cluster (C_7) exhibited a
spectrally triphasic response. BCs also differed in their temporal responses. For
example, while cluster C_2 consistently responded in a sustained manner, cluster C_3
responses were more transient and overshot the baseline between light-flashes.
Finally, diverse spectral and temporal response differences did not only exist
between BC clusters, but also within. For example, cluster C_6 switched from transient
responses during long-wavelength stimulation to sustained responses during short-
 wavelength stimulation. In some cases, such intermixing of spectral and temporal
encoding in a single functional BC-cluster could be quite complex. For example,
cluster C_21 switched from small transient On-Off responses via intermediate
amplitude transient-sustained On-responses to large amplitude sustained-only On-
responses in a wavelength-dependent manner.

Overall, in line with connectivity^{12,38} and previous functional work, both the
spectral^{19,22,23} and the temporal diversity^{19,22,23,30,39,40} of larval zebrafish BCs long
exceeded that of the cones, which at the level of presynaptic calcium were generally
sustained^{27}, and which only exist in four spectral variants (cf. Figure 1B).
Linear cone-combinations using four temporal components can account for BC responses. We next explored if and how these BC cluster-means (Figure 2B) could be explained based on cone responses\textsuperscript{27} (Figure 3, cf. Figure 1B). For this, we implemented a simple linear model (Methods) based on the following considerations.

BCs may receive cone inputs by two main, non-mutually exclusive routes: directly, via dendritic contacts onto cone-pedicles in the outer retina, and indirectly, via lateral inputs from amacrine cells in the inner retina\textsuperscript{28}. A third route, via horizontal cells, has been proposed in the case of mice\textsuperscript{41}. If such a route exists in zebrafish remains unknown.

In the outer retina, direct cone inputs are based on BC-type specific expression of glutamate receptor and/or transporter variants that are thought to be either all-sign-conserving or all-sign-inverting, but apparently never a mixture of both\textsuperscript{28,42}. Accordingly, dendritic inputs alone should only be able to produce spectral tuning functions in BCs that can be explained by same-sign cone inputs. Any BC that cannot be explained in this manner is then expected to require spectrally distinct inputs from amacrine cells. On the other hand, variations to the temporal structure of a given cones’ contribution to a BC’s response could be implemented via either route\textsuperscript{28,35,43} – that is, via a combination of dendritic and/or axonal inputs. Accordingly, we reasoned that for a linear transformation, each cone-type may feed into a functional BC-type via a unique temporal profile that represents the sum of all routes from a given cone to a given BC. In this way, our model effectively sought to explain each BC-cluster as a weighted sum of four spectral cone-tunings, but each of these four cone-inputs could have a unique temporal structure.
To capture the above considerations in a linear model, we combined the four-cone spectral tuning functions (Figure 3A, cf. Figure 1B) with four dominant temporal components extracted from BC responses: light-transient, light-sustained, dark-transient, and dark-sustained (Figure 3B, Methods). We restricted the model to capture the central ten light-stimuli (i.e. omitting the first two red-flashes and the last UV-flash) where BC-clusters generally exhibited the greatest response diversity (Figure 2).

Notably in the following paragraphs, we avoid the use of the common shorthand “On” or “Off” because in view of spectral opponency already present in cones27 a sign-conserving input to a BC is not categorically “Off”, and vice versa a sign-inverting input is not categorically “On”. Instead, we use the terms “light” and “dark” response, in reference to a response that occurs in the presence or absence of a light-stimulus, respectively. Also note that all extracted spectral tuning functions (e.g. Figure 3A) are x-inverted compared to the time-axes in recordings and reconstructions (e.g. Figure 3D,E). This was done because recordings were performed from long- to short wavelength stimuli, but spectral tuning functions are conventionally plotted from short- to long-wavelengths. Weights were scaled such that the mean of their magnitude equalled one, with weights <0.5 (“near-zero”) excluded from the summary plots for visual clarity. Full weights, including a detailed overview of each cluster, are available in Data S1.

Figures 3C-E illustrate the intermediate steps (Figure 3C,D) and final output (Figure 3E) of the model for example cluster C22. This functional BC-type was broadly tuned but switched from transient responses to long wavelength stimulation to more sustained responses at shorter wavelengths (Figure 3E, grey trace, cf. Figure 2A,B). To capture this behaviour (Figure 3E, black trace), the model drew on all four cones.
(Figure 3C), however with a particularly strong sign-conserved contribution from red-cones (Figure 3C, left). Here, the model placed a strong sign-conserving weight onto the dark-transient (Dtr) component of the red-cone (Figure 3D, left, third trace). The strength and sign of this weight is illustrated in Figure 3C (third downwards facing red bar). In addition, the model also placed weaker sign-conserving weights onto the dark-sustained (Figure 3D, left, fourth trace) and light-sustained (second trace) components, and a weak sign-inverted weight onto the dark-transient component (first trace). Summation of these four kinetic components yielded the total modelled red-cone contribution to this cluster (Figure 3D, bottom trace).

The same principle was applied across the remaining three cones, yielding a total of sixteen (four cones times four temporal components) weights per cluster (cf. Figure 3C). In the example presented, weights were mostly sign-conserving (facing downwards). However, to capture the relatively complex temporal dynamics of this cluster, which systematically overshot the baseline between flashes, the model also drew on a number of weaker sign-inverted weights (facing upwards), for example for all light-transient components.

Figure 3F illustrates mean outputs of the model for another four example clusters with diverse spectral and temporal behaviours. Of these, the spectrally bimodal but “temporally simple” response profile of C1 was well-approximated by all sign-conserving inputs from red- and UV-cones (Figure 3F, left). Similarly, the spectrally opponent behaviour of C15 could be captured by all-sign-conserving inputs from all four cones (Figure 3E, second panel). Accordingly, as expected from the cone-tunings, generating opponent responses at the level of BC terminals does not categorically require new sign-opposition in the inner retina – instead, the opponency can simply be inherited from the cones. Nevertheless, not all opponent BC
responses could be explained in this manner. For example, opponent cluster $C_{14}$ required sign-inverted inputs from red-cones but sign-conserving inputs from green-, blue- and UV-cones (Figure 3E, third panel). Finally, even the more complex spectral and temporal BC-clusters could be well-approximated by relatively simple cone-mixtures. For example, $C_{25}$ was captured by combining sign-conserved light- and dark-transient inputs from red- and blue-cones with mostly sustained and sign-inverted inputs from UV-cones (Figure 3E, rightmost).

Overall, this linear fitting procedure captured $\sim 95\%$ of the total variance across the 29 cluster means (Figure S2A, Methods). Similarly, the fits also captured $\sim 95\%$ of the temporal detail, based on comparison of the mean power spectra of the cluster means and that of the residuals (Figure S2B, Methods). The full result of this process is summarised in Figure 4, each time showing the cluster mean (grey) and reconstruction (black) alongside weight-summaries per cone following the schema illustrated in Figure 3B,C. Further detail is shown in Data S1.

Based on the traditional separation of the inner retina into “Off-” and “On-layers”$^{28}$, we may correspondingly expect mainly sign-conserving (negative) weights in “Off-stratifying” clusters $C_1$-$C_{18}$, and mainly sign-inverting (positive) weights for “On-stratifying” clusters $C_{19}$-$C_{29}$. However, this expectation was not met in several cases, for example for most of the On-stratifying clusters which nevertheless showed a general abundance of negative (“Off”) weights for red-, green- and blue-cone inputs. From here, we next explored the general rules that govern overall cone-signal integration by BCs.
The inner retina is dominated by red-cone inputs. First, we computed histograms of all weights per cone (Figure 5A) and per temporal component (Figure 5B) to determine the dominant input-motifs across the population of all BCs. This revealed that overall, the amplitudes of red-cone weights tended to be larger than those of all other cones (red absolute weights $W_R = 1.82 \pm 1.22$; $W_{G,B,U} = 0.68 \pm 0.47, 0.62 \pm 0.45, 0.87 \pm 0.88$, respectively, range in SD; $p<0.001$ for all red-combinations, Wilcoxon Rank Sum Test). This red-dominance was stable also when the four eye-regions were analysed separately ($p<0.001$ in each case). Similarly, light-response component weights tended to be larger than dark-response component weights ($W_{LT, LS, DT, DS} = 0.94 \pm 0.75, 1.73 \pm 1.20, 0.85 \pm 0.8, 0.48 \pm 0.54$, respectively Figure 5B). Here, the light-sustained response components that already dominate the cones (cf. Figure 1B) remained largest overall also in BCs ($p<0.001$ for all Light$\text{sus}$-combinations, Wilcoxon Rank Sum Test).

Red-, green- and blue-cone weights co-vary independent of UV-cone weights. Next, we explored the weight relationships between the four cone types across clusters. In general, a strong correlation between weights attributed to any two cone types would suggest that inputs from these cones tend to be pooled, for example by the dendrites of individual BCs contacting both cone-types. In contrast, a low correlation or even anticorrelation between cone-weights could indicate the presence of cone-opponency.

Across clusters, we found that red-cone weights strongly correlated the weights of both green- ($\rho = 0.73$; 95% confidence intervals (CI) 0.49/0.86, Figure 5C) and blue-cones ($\rho = 0.87$, CI 0.74/0.94, Figure 5D; green vs. blue: $\rho = 0.89$; CI 0.77/0.95, cf.
Figure S3A). The tight association between red-, green- and blue-cone weights extended across both the all-sign inverting (bottom left) and the all-sign-conserving (top right) quadrants and comprised few exceptions in the two remaining quadrants that would indicate cone-opponency. Accordingly, zebrafish BCs did not tend to differentially combine inputs from red-, green- or blue-cones of either polarity to set up potentially new opponent-axes.

In contrast, red-cone weights were uncorrelated with UV-cone weights ($\rho = -0.21$, CI -0.55/0.14, Figure 5E, green sc. UV: $\rho = -0.04$, CI -0.40/0.34; blue vs. UV: $\rho = -0.34$, CI -0.63/0.03, see Figure S3B,C), with many clusters scattering across the two sign-opponent quadrants (i.e. top left, bottom right). Accordingly, reconstructing a substantial fraction of BC clusters required opposite sign inputs from red-/green/blue-versus UV-cones, suggestive of a newly set-up form of spectral opponency in the inner retina. Interestingly, in some cases, a cluster could exhibit small “Off” responses in the UV range despite using sign-inverting weights for UV-cones (e.g. C23 – best seen in Data S1). This was possible because all cones respond to UV-light to some extent (Fig. 1B). In some cases, the sum of inferred red-, green, and blue-cone inputs could then outweigh inferred UV-cone inputs. We next explored the spectral tuning of BC-clusters in further detail.

BC spectral responses fall into three opponent, and three non-opponent groups. The complex interplay of temporal and spectral structure in BC-responses (Figure 2) meant that their spectral tuning functions could not easily be extracted directly from the BC-cluster means, for example by means of taking the area under the curve in response to each flash of light. Instead, we estimated their tuning
functions based on their fitted cone-weights (cf. Figure 4). To this end, for each
cluster we summed sixteen cone-tuning functions (based on Figure 3A), each scaled
by the cluster’s associated sixteen weights (i.e. red-L_tr + red-L_{sus.} + red-D_tr. and so
on). This summarised each cluster’s ‘bulk’ response in a single spectral tuning
function that gave equal weight to each of the four temporal components (Figure 5F-
K). By this measure, 18 of the 29 BC-clusters were non-opponent (62%, Figure 5F-
H) and 11 were opponent (38%, Figure 5I-K). Here, opponency was defined as any
tuning function that crossed and overshot zero at least once with an amplitude of at
least 10% compared to that of the opposite (dominant) polarity peak response.

Non-opponent clusters (‘closed’ symbols, cf. Figure 4A) approximately adhered to	hree major groups: spectrally broad (three On- and eight Off-clusters, Figure 5F),
approximately UV-cone-like (one On- and four Off-clusters, Figure 5G), and
approximately red-cone-like (two Off-clusters, Figure 5H). Similarly, opponent
clusters (‘open’ symbols) fell into three major groups based on the spectral positions
of their zero crossings: Two green-cone-like clusters (both short_{off}/long_{on}, crossing at
520 and 536 nm, Figure 5I), three blue-cone-like clusters (two short_{off}/long_{on}
crossing at 497 and 499 nm, plus the single triphasic C_7 with a dominant
short_{on}/long_{off} zero crossing at 490 nm, Figure 5J), and six UV-cone versus red-/
green-/blue-cone opponent clusters (henceforth: UV:R/G/B, five short_{on}/long_{off}:
crossing at 416, 425, 428, 435, 448 nm, one short_{off}/long_{on} crossing at 438 nm,
Figure 5K). In comparison, green- and blue-cone zero-crossings, respectively
(Figure 5L, from Ref27) occurred at ~523 and ~483 nm, while red- and UV-cones,
respectively, approached zero between ~425 and 450 nm (Figure 5I-N, shadings).

The tight correspondence between opponent BC-clusters (Figure 5I-K) and cone-
tunings (Figure 5L) was further illustrated by the histogram of BC-zero-crossings that
also incorporated relative abundances of ROIs contributing to each cluster (Figure 5M). The histogram showed three clear peaks that were well-aligned to the three spectral axes set-up in the cones (shadings). Further, the histogram also retained its overall shape when the four temporal components underpinning each cluster were considered individually (Figure 5N). As a control, this trimodal structure disappeared when component-weights were iteratively randomised (Figure S3D), or when temporal-components were randomly shuffled between cones (Figure S3E), suggesting that the measured BC tunings emerged from non-random effective cone-inputs. In support, and despite appreciable diversity, the spectral tuning functions of the four temporal components that contributed to a given cluster tended to be positively correlated among both opponent and non-opponent clusters (Figure S3F-I).

Remarkably therefore, it appears that by and large, BCs tended to retain many of the dominant spectral properties of the cones rather than build fundamentally new spectral axes – all despite integrating across multiple cone types and presumably diverse inputs from spectrally complex ACs\(^\text{23}\). The only two notable deviations from this observation were a highly stereotypical spectral broadening in 11 clusters (Figure 5F), which may be linked to outer retinal cone-pooling\(^\text{12}\), and, strikingly, the emergence of six strongly UV:R/G/B opponent clusters (Figure 5K).

**UV-cone, but not red-/green-/blue-cone weights follow traditional IPL On-Off lamination.** Finally, we asked where the inferred new form of UV:R/G/B opponency might be set-up in the inner retina (Figure 6). To this end, we combined the cone-weight data (Figure 4) with information about each BC-terminal’s stratification depth.
within the inner plexiform layer (IPL) (Figure 3C). In general, the IPL of all vertebrates studied to date is dominated by “Off-circuits” in the upper strata, adjacent to the somata of BCs and most amacrine cells, and by “On-circuits” in the lower strata, adjacent to the somata of retinal ganglion cells\textsuperscript{28}. Accordingly, light-components L_{tr} and L_{sus} are expected to mostly exhibit sign-conserving weights in the upper strata, and mostly sign-inverting weights in the lower strata (Figure 6A). Dark components D_{tr} and D_{sus} are expected to exhibit the reverse distribution (Figure 6B).

This textbook expectation, here graphically indicated by dashed lines, was indeed approximately met when considering dark-components (Figure 6B - note that UV-dark component weights were generally small and not further considered) and for light-components of UV-cones (Figure 6A, bottom panel). Similarly, this classical IPL organisation was also met by red-, green- and blue-cone weights for the upper two-thirds of the IPL, which included the traditional Off-layer, and the upper part of the traditional On-layer (Figure 6A, top three panels). However, specifically for red-, green-and blue-cones, the lower third of the traditional On-layer was dominated by weights of the “wrong” polarity (Figure 6A, top three panels). In agreement, most UV:R/G/B opponent clusters stratified in this lower third of the IPL (Figures 3C,4). Together, this suggests that several of these UV:R/G/B clusters are derived from sign-reversed red-/green-/blue-cone inputs onto “native” UV-On BCs, for example by way of amacrine cells.
We have shown that the substantial spectral and temporal diversity of larval zebrafish BCs (Figures 1,2, cf. Refs^{19,30}) can be well-captured by a linear combination of inputs from the four spectral cone-types (Figure 3,4). This in turn allowed us to explore the major functional connectivity rules that govern spectral and temporal widefield signal integration by BCs: We find that red-cones overall provide the dominant input to BCs, often complemented by weaker but same-sign inputs from green- and blue-cones (Figure 5A,C,D). Likely as one consequence, BC pathways do not generally set-up new axes of spectral opponency in the mid- to long-wavelength range. Rather, they mostly either conserve and diversify the two major opponent motifs already present in the cones (Figure 5I,J) or establish non-opponent circuits (Figure 5F-H). In contrast, inner retinal UV-cone pathways appear to be organised essentially independently to those of red-, green- and blue-cones (Figure 5E). This leads to the consolidation of a third axis of spectral opponency, contrasting long- and mid-wavelength signals against UV (Figure 5K). This third axis appears to mainly stem from a systematic polarity reversal of inputs from red-, green- and blue-cones onto ‘natively-UV-On’ BCs in the lower IPL (Figure 6A).

Building spectrally opponent BCs. Because spectral opponency is a prominent feature in larval zebrafish cones^{27}, BCs may inherit this property rather than set-up new opponent spectral axes by way of ACs. Indeed, the opponency observed in BC cluster C_{15} could be explained based on weighted but all-sign-conserving inputs from all four cones (Figure 4). However, the full picture may be more complex. For example, like C_{15}, cluster C_{14} was also opponent, albeit with a stronger long-
wavelength response, and in this case the model used weakly sign-inverted red-
cone weights alongside sign-conserved green- and blue-cone weights. In fact, most
UV:R/G/B opponent clusters (e.g. C_{25-29}) required opposition of long versus short-
wavelength cone inputs in the inner retina. This hints that inner retinal circuits may
generally use a “mix-and-match” strategy to achieve diverse spectral responses by
any available route, rather than strictly adhering to any one strategy. This notion is
also tentatively supported by the presence of spectrally diverse amacrine cell circuits
in adult zebrafish^{23}. More generally, it perhaps remains puzzling how the complex
interplay of cone pooling in the outer retina with AC inputs in the inner retina, across
29 highly diverse functional-BC-types which presumably express diverse receptors
and ion channels^{28}, can ultimately be summarised in an functional wiring logic that
for the most part simply sums all four cones, or ‘at best’ opposes a red-/green-/blue-
system against UV. Resolving this conceptual conflict will likely require targeted
circuit manipulations, for example by comparing BC spectral tunings in the presence
and absence of amacrine cell inputs, or after targeted cone-type ablations.

Beyond ‘classical’ opponency, several clusters – both opponent and non-opponent –
in addition encoded a notable mixture of spectral and temporal information.
Interestingly, several of these clusters appeared to be concentrated around the
centre of the IPL (e.g. C_{20-25}, Figure 2B,C) – a region which also in mammals has
been associated with both transient and sustained processing^{35,44–46}. In zebrafish, a
mixed time-colour code was previously described for the downstream retinal
ganglion cells^{22}, which now raises the question to what extent ganglion cells may
inherit this property from BCs. Moreover, if and how such information can be
differentially read out by downstream circuits and used to inform behaviour remains
unknown.
Three axes of spectral opponency. In principle, the four spectral cone types of larval zebrafish could be functionally wired to for tetrachromatic vision. This would require that all four cone types contribute independently to colour vision. Theory predicts that efficient coding of colour should be based on four channels, an achromatic channel with no zero-crossings on the spectral axis, and three chromatic opponent channels with one, two and three zero-crossings respectively\textsuperscript{5,47}. However, such a coding strategy is not essential as demonstrated by the trichromatic visual system of many old-world monkeys which is based on two axes of opponency (“blue-yellow” and “red-green”), each with a single zero crossing. In the present study, we find that among zebrafish BCs, three zero-crossings predominate (Figure 5M,N, Figure 7A). Here, the single BC cluster with two zero-crossings (C\textsubscript{7}) did not set-up any notable additional spectral crossings either, but instead crossed once in the ‘blue-cone position’, and once again near the ‘UV-red opponent position’ (Figure 5K). Nevertheless, our findings support the notion that at least at the level of BCs, and under the stimulus conditions used in this study, the zebrafish visual system is capable of supporting tetrachromatic colour vision, as observed behaviourally in goldfish\textsuperscript{48}. If and how the larval zebrafish BCs’ axes are preserved, diversified, or even lost in downstream circuits will be important to explore in the future. In this regard, both retinal ganglion cells\textsuperscript{21,22} and brain circuits\textsuperscript{21,49} do carry diverse spectral signals, however beyond a global overview\textsuperscript{29} the nature and distribution of their spectral zero-crossings remain largely unexplored.

Links with mammalian SWS1:LWS opponency. Of the three spectral axes that dominate the zebrafish inner retina (Figure 5I-N, 7A), those functionally linked with green- (RH2) and blue-cone (SWS2) circuits are unlikely to have a direct counterpart
in mammals where these cones-type are lost\textsuperscript{1,9}. However, the third axis, formed by functional opposition of UV-cone circuits against red-/green-/blue-cone circuits, may relate to one or multiple of the well-studied mammalian SWS1:LWS opponent circuits\textsuperscript{50,51} (Figure 7B).

Despite substantial spectral variation amongst both SWS1 and LWS cone-types across species, mammals usually oppose the signals from SWS1-cones with those of LWS-cones at a retinal circuit level\textsuperscript{4,6,52–56}. For example, in the primate outer retina, SWS1-cones exhibit horizontal-cell mediated spectral opponency to LWS signals\textsuperscript{57}. Likewise, in the inner retina signals from a highly conserved SWS1-exclusive On-BCs are combined with those of LWS-biased Off-circuits in most if not all mammals that have been studies at this level\textsuperscript{36,51,58,59}. Further such circuit motifs can involve diverse but specific types of amacrine and/or retinal ganglion cells\textsuperscript{4,54,60}.

Several of these mammalian motifs may have a direct counterpart in zebrafish. For example, like primate SWS1-cones, also zebrafish SWS1-cones exhibit weak but significant long-wavelength opponency that is mediated by horizontal cells\textsuperscript{27}. Beyond this possible outer retinal connection, the inferred UV:R/G/B organisation in zebrafish BCs (Figures 5E,K, 6) is reminiscent of mammalian circuits associated with SWS1-BCs.

First, as in most mammals\textsuperscript{52}, SWS1\textsubscript{On}:LWS\textsubscript{Off} signals numerically dominate in zebrafish compared to SWS1\textsubscript{Off}:LWS\textsubscript{On} signals. Second, zebrafish SWS1:LWS opponent signals are predominately found in the lower-most (GCL-adjacent) fraction of the IPL (Figures 3, 6), the same place where mammalian SWS1-On BCs stratify\textsuperscript{36}. Third, many zebrafish SWS1\textsubscript{On}:LWS\textsubscript{Off} signals occurred ventro-temporally (Figure 3D), the retinal region which in mice exhibits the highest density of type-9 BCs\textsuperscript{61},...
their only SWS1-exclusive BC type\textsuperscript{36,58}. While zebrafish are not known to possess an SWS1-exclusive BC\textsuperscript{12}, they do possess several anatomical BC types that contact SWS1-cones alongside either one or both of SWS2- (blue) and RH2-cones (green)\textsuperscript{9,12}. Such BCs may conceivably become SWS1-exclusive types upon the loss of RH2 and SWS2 cones in early mammalian ancestors.

However, not everything supports a direct correspondence between mammalian and zebrafish SWS1:LWS circuits. For example, in contrast to BCs, among the dendrites of the zebrafish retinal ganglion cells, most UV-opponent signals occur above the IPL midline, near the anatomical border between the traditional On- and Off-layers\textsuperscript{22}. Nevertheless, this is approximately in line with the IPL position where several of the well-studied primate SWS1:LWS ganglion cells receive LWS-biased Off-inputs\textsuperscript{62}, hinting that similar ganglion cell motifs might also exist in zebrafish. Certainly, zebrafish do possess a number of anatomical retinal ganglion cell types\textsuperscript{22,63} that display similar stratification patterns compared to those that carry SWS1:LWS opponent signals in diverse mammals\textsuperscript{51,54}.

A summary of the above argument, showcasing possible links between retinal circuits for colour vision in cone-tetrachromatic species such as zebrafish, to those of most non-primate mammals and of old-world monkeys including humans, is suggested in Figure 7A-C. In the future it will be important to explore if and how mammalian circuits such as the ones carrying SWS1:LWS signals can be more directly linked with those found in zebrafish, for example by leveraging molecular markers across potentially homologous types of neurons\textsuperscript{37,64,65}. 
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Author contributions

Conceptualization, PB, TY, TB; Methodology, PB, TY, FKJ; Investigation, PB, TY; Data Curation, PB, TB; Writing – Original Draft, TB; Writing – Review & Editing, TB, TY, FKJ; Visualization, TB, PB; Supervision, TB; Project Administration, TB; Funding Acquisition, TB.

Declaration of Interests

The authors declare no competing interests.
**Figure 1 | Measuring high-spectral resolution tuning curves in zebrafish bipolar cells.**

A, Schematic of the larval zebrafish retina, with cone-terminals in the outer retina and bipolar cell (BC-) terminals in the inner retina highlighted. B, Mean calcium-responses of red- green-, blue- and UV-cone terminals to a series of 13 spectrally distinct widefield flashes of light as indicated (data from Ref27). Note that for clarity the response to a 14th “low-power-control” UV-LED was graphically removed compared to the original publication. C-F, Illustration of recording strategy for BC-terminals in the inner plexiform layer (IPL), and exemplary results. An optical tri-plane approach (C, top) was used to simultaneously record from three planes of larval zebrafish BC-terminals expressing SyGCaMP6f by way of two-photon imaging coupled with remote focussing (Methods). From here, we automatically placed regions of interest (ROIs) and detected the boundaries of the IPL (D, Methods). Time traces from all ROIs in a recording plane were z-scored and averaged across 3-5 response repeats of the full stimulus sequence (E). Example traces from individuals ROIs (F) are shown as individual repeats (grey) and averages across repeats (black). Zebrafish larva schematic (A) by Lizzy Griffith. See also Figure S1.

**Figure 2 | Clustering into 29 functional BC-types.**

A-D, Overview of the result from unsupervised clustering of all BC-data recorded as shown in Figure 1 that passed a minimum quality index (QI>0.4, Methods). For each cluster, shown are the individual BC-mean responses as heatmaps (A), the corresponding cluster means and SD shadings, with approximate baseline indicated in dashed (B), distribution of ROI positions in the IPL (C) and each cluster’s distribution across the four recording
regions within the eye (D, from left: acute zone, dorsal, nasal, ventral). Histograms in (C) are area-normalised by cluster, and in (D) by recording region. Clusters are ordered by their average anatomical position in the IPL, starting from the border with the inner nuclear layer (cf. C). The coloured symbols indicate the overall spectral group as assigned later (cf. Figure 5F-K).

Figure 3 | Reconstructing bipolar cell responses from cones. A-E, Summary of the reconstruction strategy for example cluster C22 (for details see Methods). Each BC-cluster reconstruction is based on the linear combination of the spectral tunings functions of the four cone-types (A, from Ref27) with four stereotyped temporal components associated with individual light flashes (B), yielding 4 X 4 = 16 weights (C). Weights are shown in blocks of temporal component weights (from left: Light-transient, Light-sustained, Dark-transient, Dark-sustained) associated with each cone (indicated by the corresponding colours). Bars above zero indicate sign-inverted (“On-”) weights, while bars below zero indicate sign-conserved (“Off-”) weights. The corresponding full expansion of this reconstruction is shown in (D). Individual combination of each cone’s tuning function (A) with each temporal component (B), scaled by their corresponding weight (C), yields sixteen “sub-traces” (D, upper four traces in each of the four panels, labelled Ltr, Lsus, Dtr, Dsus). Summation of each cone’s four sub-traces yields that cone’s total contribution to the cluster (D, bottom traces, labelled “sum”). Finally, summation of the four cone-totals yields the full reconstruction (E, black trace), shown superimposed on the target cluster mean (grey). F, as A-E, but showing only the weights (top) cone-totals (middle) and full reconstructions (bottom) for another four example clusters (from left:
Further detail on reconstructions is shown in Figure S2, and all cluster’s individual results are detailed in Data S1.

**Figure 4 – A functional overview of cone-bipolar cell mappings.** A,B, Overview of all BC-cluster means (A, grey traces, cf. Figure 2B) and their full reconstructions based on the strategy detailed in Figure 3 (black traces). Associated weights are shown in (B). For clarity, “near-zero” weights (abs(w)<0.5) are omitted. Full weights are shown in Data S1. Note that based on outer retinal inputs only, weights are generally expected to be sign-conserving for clusters in the traditional “Off” layer (C1-C18), and sign-inverting in the anatomical “On” layer (C19-C29), as indicated on the right. The round symbols plotted next to each cluster (A) denote their allocated spectral group, as detailed in Figure 5F-K and associated text.

**Figure 5 – Major trends in cone-weights and spectral tunings.** A,B, Histograms of all weights associated with inputs to each of the four cones across all clusters, independent of temporal-component types (A), and correspondingly histograms of all weights associated with temporal components, independent of cone-type (B). “Near-zero” weights (abs(w)<0.5) are graphically de-emphasised for clarity. All weights contributed equally to these histograms, independent of the size of their corresponding cluster. C-E, Scatterplots of all clusters’ weights associated with each cone plotted against each other as indicated. Large symbols denote the mean weight associated with each cone and cluster across all four temporal components (i.e. one symbol per cluster), while small symbols denote each weight individually (i.e. four symbols per cluster, corresponding to Ltr, Lsus, Dtr, Dsus). The remaining three
possible cone-correspondences (G:B, G:U, B:U) are shown in Figure S3A-C. 

Peak-normalised ‘bulk’ spectral tuning functions of all 29 clusters, grouped into six categories as indicated. The strength of each line indicates the numerical abundance of ROIs belonging to each cluster (darker shading = larger number of ROIs; exact number of ROIs contributing to each cluster are listed in Data S1). As appropriate, spectral tuning functions of cones (cf. L) are shaded into the background, as appropriate (G,H, thick coloured traces) to illustrate the close spectral correspondences of associated cones and BCs. Similarly, for three spectrally opponent groups (I-K), the approximate positions of the corresponding cone’s zero crossings are indicated with a vertical shaded line (cf. L). L, Cones’ spectral tuning functions, with approximate zero-crossings (blue-/green-cones) and zero-positions (red-/UV-cones) graphically indicated. M,N, Histograms of zero-crossings across all BC-clusters, incorporating the abundance of ROIs belonging to each cluster. Shown are crossings of ‘bulk’ spectral tunings functions (M, cf. F-H), and of spectral tuning functions that were computed for each temporal component individually, as indicated (see also Figure S3F-I, and Data S1). Note the three prominent peaks of zero-crossing positions, approximately aligned with the zero-positions/crossings of the cones. These peaks largely disappeared when time-components were fully randomised (Figure S3D) or randomly permuted across cones (Figure S3E).

**Figure 6 – Cone-weight distribution across the inner plexiform layer.** A,B, Two-dimensional histograms of weights (x-axes) associated with each cone resolved by IPL position (y-axes). Brighter colours denote increased abundance. For simplicity, the weights associated with the light (Ltr, Lsus) and dark-components (Dtr, Dsus), are combined in panels A and B, respectively. Moreover, near-zero weights are not
shown (central white bar in all panels). The thick white dotted lines indicate approximate expected distribution of weights based on traditional “On-Off” lamination of the inner retina. By each panel’s side, instances where this expectation is violated are highlighted as “polarity violation”.

**Figure 7 – Possible links across vertebrate retinal colour circuits.** A-C, Conceptual summary schematics of retinal circuits for colour vision in zebrafish (A), dichromatic mammals such as many rodents (B) and some trichromatic old-world monkeys such as humans (C). The coloured ‘graphs’ indicate approximate spectral tuning functions of retinal neurons in a given layer, as indicated.

**STAR METHODS**

**RESOURCE AVAILABILITY**

**Lead Contact**

Further information and requests for resources and reagents should be directed to and will be fulfilled by the Lead Contact, Tom Baden (t.baden@sussex.ac.uk).

**Data and Code Availability.** Pre-processed functional 2-photon imaging data and associated summary statistics is freely available on DataDryad under https://doi.org/10.5061/dryad.wstqiq2n566 and via the relevant links on http://www.badenlab.org/resources and http://www.retinal-functomics.net. See also Data S1 for a graphical summary of key aspects pertaining to each BC cluster.
**Materials Availability.** The transgenic line Tg(1.8ctbp2:SyGCaMP7bf) used in this study is available upon request to the lead author.

**EXPERIMENTAL MODEL AND SUBJECT DETAILS**

**Animals.** All procedures were performed in accordance with the UK Animals (Scientific Procedures) act 1986 and approved by the animal welfare committee of the University of Sussex. Animals were housed under a standard 14:10 day/night rhythm and fed three times a day. Animals were grown in 0.1 mM 1-phenyl-2-thiourea (Sigma, P7629) from 1 dpf to prevent melanogenesis. For all experiments, we used 6-7 days post fertilization (dpf) zebrafish (Danio rerio) larvae.

Tg(1.8ctbp2:SyGCaMP7bf) line was generated by injecting pBH-1.8ctbp2-SyjGCaMP7b-pA plasmid into single-cell stage eggs. Injected fish were out-crossed with wild-type fish to screen for founders. Positive progenies were raised to establish transgenic lines. The plasmid was made using the Gateway system (ThermoFisher, 12538120) with combinations of entry and destination plasmids as follows: pBH and p5E-1.8ctbp, pME-SyjGCaMP7b, p3E-pA. Plasmid p5E-1.8ctbp was generated by inserting a polymerase chain reaction (PCR)-amplified -1.8ctbp fragment into p5E plasmid and respectively. Plasmid pME-SyjGCaMP7b was generated by replacing GCaMP6f fragment with PCR-amplified jGCaMP7b in pME-SyGCaMP6f plasmid.

For 2-photon *in-vivo* imaging, zebrafish larvae were immobilised in 2% low melting point agarose (Fisher Scientific, BP1360-100), placed on a glass coverslip and submerged in fish water. Eye movements were prevented by injection of α-
bungarotoxin (1 nL of 2 mg/ml; Tocris, Cat: 2133) into the ocular muscles behind the eye.

METHOD DETAILS

**Light Stimulation.** With fish mounted on their side with one eye facing upwards towards the objective, light stimulation was delivered as full-field flashes from a spectrally broad liquid waveguide with a low numerical aperture (NA 0.59, Newport), positioned next to the objective at ~45°, as described previously\(^\text{27}\). To image different regions in the eye, the fish was rotated each time to best illuminate the relevant patch of photoreceptors given this stimulator-geometry. The other end of the waveguide was positioned behind a collimator-focussing lens complex (Thorlabs, ACL25416U-A, LD4103) which collected the light from a diffraction grating that was illuminated by 13 spectrally distinct light-emitting diodes (LEDs, details below). After mounting but before systematic light stimulation, fish were exposed to at least 5 minutes of “spectral noise” (each LED independently flickering in a random sequence) to light-adapt the eye.

An Arduino Due (Arduino) and LED driver (Adafruit TCL5947) were used to control and drive the LEDs, respectively. Each LED could be individually controlled, with brightness defined via 12-bit depth pulse-width-modulation (PWM). To time-separate scanning and stimulating epochs, a global “blanking” signal was used to switch off all LEDs during 2P scanning but enable them during the retrace, at line-rate of 1 kHz (see also Refs\(^\text{71,72}\)). The stimulator code is available at [https://github.com/BadenLab/HyperspectralStimulator](https://github.com/BadenLab/HyperspectralStimulator).
LEDs used were: Multicomp Pro: MCL053RHC, Newark: C503B-RAN-CZ0C0AA1, Roithner: B5-435-30S, Broadcom: HLMP-EL1G-130DD, Roithner: LED-545-01, TT Electronics: OVLGC0C6B9, Roithner: LED-490-06, Newark: SSL-LX5093USBC, Roithner: LED450-03, VL430-5-1, LED405-03V, VL380-5-15, XSL-360-5E. Effective LED peak spectra as measured at the sample plane were, respectively (in nm): 655, 635, 622, 592, 550, 516, 464, 448, 427, 407, 381, 360 nm. Their maximal power outputs were, respectively (in µW): 1.31, 1.06, 0.96, 0.62, 1.26, 3.43, 1.47, 0.44, 3.67, 0.91, 0.24, 0.23, 0.20. From here, the first ten LEDs (655 – 427 nm) were adjusted to 0.44 µW, while the three UV-range LEDs were set to a reduced power of 0.2 µW. This relative power reduction in the UV-range was used as a compromise between presenting similar power stimulation across all LEDs, while at the same time ameliorating response-saturation in the UV-range as a result of the UV-cones’ disproportionately high light sensitivity. The same strategy was used previously to record from cones.

2-photon calcium imaging. All 2-photon (2P) imaging was performed on a MOM-type 2P microscope (designed by W. Denk, MPI, Martinsried; purchased through Sutter Instruments/Science Products) equipped with a mode-locked Ti:Sapphire laser (Chameleon Vision-S, Coherent) tuned to 927 nm for SyGCaMP7b imaging. Notably, like all calcium imaging, the biosensor exhibits non-instantaneous binding and unbinding kinetics, which in effect low-pass filters the “real” calcium signals in BCs. We used one fluorescence detection channel (F48x573, AHF/Chroma), and a water immersion objective (W Plan-Apochromat 20x/1,0 DIC M27, Zeiss). For image acquisition, we used custom-written software (ScanM, by M. Mueller, MPI,
All data was collected using a quasi-simultaneous triplane approach by leveraging an electrically tunable lens (ETL, EL-16-40-TC-20D, Optotune) positioned prior to the scan-mirrors. Rapid axial-jumps of ~15 µm between scan planes (ETL settling time of <2 ms\textsuperscript{32}) were enabled by using a non-telecentric (nTC) optical configuration (nTC\textsubscript{1}, 1.2 mm – see Ref\textsuperscript{32}). This nTC optical setup is described in detail elsewhere\textsuperscript{32}. All recordings were taken at 128 x 64 pixels/plane at 3 planes (5.2 Hz effective “volume” rate at 1 ms per scan line).

**Pre-processing of 2-photon data, IPL detection and ROI placement.** Raw fluorescence stacks were exported into a Python 3 (Anaconda) environment. The data were de-interleaved and separated into the three recording planes. Next, the data were linearly detrended, linearly interpolated to 42 Hz, and aligned in time. The anatomical borders of the inner plexiform layers were automatically detected by first median-smoothing the time standard deviation images with a Gaussian kernel size of 3 pixels. From here, every pixel above the 35% per-image amplitude threshold was registered as IPL. This automated procedure was made possible by the fact that GCaMP6f expression was restricted to the presynaptic terminals of BCs, which also defined the anatomical borders of the IPL.

To place regions of interest (ROI), a quality index (QI) as described previously\textsuperscript{34} was calculated for each pixel. In short, the QI measures the ratio of variance shared between stimulus repetitions and within a single stimulus repetition. The larger the QI, the more variance in the trace is due to the presented stimulus:

$$QI = \frac{VarC_{rr}}{VarC_{rr}}$$
where $C$ is the $T$ by $R$ response matrix (time samples by stimulus repetitions) and $x$ and $\text{Var}[x]$ denote the mean and variance across the indicated dimension, respectively. $QI$ ranges from 0 (perfectly random) to 1 (all stimulus repetition responses are identical). This yielded “$QI$-images” that indicated where in a scan BC-responses were located. From here, ROIs were automatically placed using custom Python scikit-image scripts\textsuperscript{73}. In brief, $QI$-images were adaptively thresholded using kernel size 5 pixels which helped accentuate responsive image structures that were approximately BC-terminal-sized (in our scan configuration, most BC-terminals were \~5 pixels in diameter – cf. Figure 1D). The resulting binary images were distance-transformed and shrunk. The contours of the remaining groups of pixels were recorded and filled, and the highlighted pixels were used as ROI coordinates. This yielded ROI sizes of $1.36\pm0.17$ $\mu m^2$ (mean$\pm$SD), which is in line with anatomical sizes of BC terminals in larval zebrafish\textsuperscript{74}. While it remains possible that a minority of ROIs over- or under-split terminals, this possible limitation was judged to be minimal based on manual inspections. The IPL position of each ROI was defined as the relative position of the centre-of-mass of the filled ROI contour to the nearest inner and outer borders of the IPL.

ROI traces were converted to z-scores. For this, a 5 s portion of the trace preceding stimulus presentation was drawn and defined as baseline. The standard deviation of this baseline fluorescence signal was calculated and used to z-score the remainder of the trace. Finally, $QIs$ as described above for each pixel were also calculated for each ROI. In line with how we previously processed the cones\textsuperscript{27}, ROIs with $QI<0.4$ were excluded from further analysis. $n = 6,125$ ROIs passed this quality criterion (72 triplane scans from 7 fish).
**Clustering of BCs.** To identify structure amongst the BC-dataset, trial-averaged ROI traces were PCA-transformed and clustered as described previously (e.g. Refs^{19,34}). In brief, we used the first 48 principal components, which accounted for 82% of total variance. Of these, components that near-exclusively carried high-frequency content which is likely linked to noise were discarded. The transformed time-traces were clustered using the scikit-learn (Python 3, Anaconda) implementation of the Gaussian Mixture Models algorithm. The number of clusters (29) was determined using the Bayesian information criterion (BIC). However, the BIC curve notably flattened above ~20 clusters, suggesting that a range of solutions would be similarly plausible. Clusters were judged as stable over repeated clustering runs starting from different random seeds, in the sense that they always picked up several broadband and UV:R/G/B response types, followed by a smaller number of “cone-like” ones (cf. Figure. 5).

**Reconstruction of BC responses from cones.**

To reconstruct each BC-mean response into constituent spectral and temporal components, we combined the average spectral tuning curve of each of the four cone-types (from Ref^{27}) with four temporal components associated with a given light response (i.e. 1.5 s On, 1.5 s Off). The four temporal components used, obtained by non-negative matrix factorisation across all light responses and cluster means, resembled light-transient, light-sustained, dark-transient, and dark-sustained temporal profiles (Figure 3B). Next, each ROI’s trial averaged trace was decomposed into a corresponding 4 by 10 array (four temporal components X 10 LEDs). Here, we restricted the reconstruction to the central 10 LEDs that generally elicited the greatest variance across BCs. This also avoided using responses to the
shortest wavelength LED which may have driven saturating responses in UV-cones (UV-cones are more light-sensitive than the other cones). Moreover, it avoided using the two longest-wavelength LEDs where responses were comparatively weak and thus noisy. This yielded four spectral tuning curves per ROI (i.e. light-transient x 10 LEDs, light-sustained x 10 LEDs and so on), which were then linearly interpolated to the range of 360 - 610 nm to conform with the cone data format. The BC tuning curves were then modelled as linear combinations of the cone tuning curves with a lasso regulariser, which yielded four cone weights X four response bases per BC-trace. For simplicity, we henceforth used the ROI-averaged weights within a cluster for further processing, but each ROI's individual weights are available to download from DataDryad[^66].

To assess reconstruction quality (Figure S2), reconstructed data was subtracted from the original ROI-means to yield residuals. From here, we compared original data, reconstructions, and residuals by two metrics: variance explained across all clusters, and temporal power explained. To determine the fraction of variance explained by the reconstructions, we first computed the total variance across all clusters for each time-point. The result of this process, plotted beneath each corresponding heatmap (Figure S2A), showed similar time-variance profiles across cluster means and their reconstructions (panels 1 and 2), but very little remaining signal for the residuals (panel 3). From here, we computed the area under the curve for each variance-trace and normalised each to the result from the original cluster means. By this metric, cluster reconstructions captured 94.0% of the original variance, while residuals carried 5.1%.
To determine the extent to which temporal structure was captured, we used a similar approach to the one above, however in this case based on a magnitude-squared Fourier Transform of each time-trace (Figure S2B), limiting the result between 0.16 and 2 Hz which captured the bulk of physiologically meaningful temporal components given the optical imaging approach used (i.e. lower-frequency components would mainly arise from imperfect detrending, while higher-frequency components would exceed the Nyquist recording limit, and further be limited by the kinetics of GCaMP7b. From here, we computed the average of all clusters’ Fourier transforms (plotted beneath each panel) and again computed the fraction of this signal captured by the reconstruction (103.8%) and residuals (3.8%). Notably, while this metric was mainly informative about low frequency components which dominated all signals, also higher frequency components were generally well captured, as visible in the individual heatmaps.

QUANTIFICATION AND STATISTICAL ANALYSIS

Statistics. No statistical methods were used to predetermine sample size. Owing to the exploratory nature of our study, we did not use randomization or blinding. To compare weight amplitude distributions (Figure 5A,B) we used the paired Wilcoxon Rank Sum Test, taking paired components as the input (i.e. comparing red-light-transient versus green-light-transient, and so on). To assess weight correlations between cones (Figure 5C-E, Figure S2), we in each case list the Pearson correlation coefficient $\rho$ and 95% confidence intervals (CI) based on the mean weights per cluster. Individual temporal weights were not considered in this analysis. All statistical analysis was performed in Python 3 (Anaconda) and/or Igor Pro 6 (Wavemetrics).
**Data S1 | Detailed Summary for each Cluster. Related to Figure 4.** For each of the 29 clusters as indicated (1 cluster per page), overview of key response aspects and analysis as shown across the main figures. Upper row, from top: Stimulus sequence, heatmap of the response-mean of all ROIs assigned to the cluster (as in Fig. 2A), cluster mean±SD (grey, as Fig. 2B) and reconstruction (black, as Fig. 4A). Lower row, from top/left: Allocated cone-weights (as Fig. 4B), here with SD error bars across each individual ROI that contributes to a cluster, and bottom left: distribution of ROIs across the IPL and eye (as Fig. 2C,D). Bottom right: Spectral tuning functions extracted from cone weights for each temporal component as indicated (thin lines, as Fig. S3F,G) and bulk tuning function based on the combination of all temporal components (thick line, as Fig. 5F-K).

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