# **Distinct Synaptic Transfer Functions in Same-Type Photoreceptors**

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5 **SUMMARY.** Many sensory systems use ribbon-type synapses to transmit their signals to 6 downstream circuits. The properties of this synaptic transfer fundamentally dictate which 7 aspects in the original stimulus will be accentuated or suppressed, thereby partially defining the detection limits of the circuit. Accordingly, sensory neurons have evolved a wide variety of 8 9 ribbon geometries and vesicle pool properties to best support their diverse functional requirements. However, the need for diverse synaptic functions does not only arise across 10 neuron types, but also within. Here we show that UV-cones, a single type of photoreceptor of 11 the larval zebrafish eye, exhibit striking differences in their synaptic ultrastructure and 12 13 consequent calcium to glutamate transfer function depending on their location in the eye. We 14 arrive at this conclusion by combining serial section electron microscopy and simultaneous "dual-colour" 2-photon imaging of calcium and glutamate signals from the same synapse in 15 vivo. We further use the functional dataset to fit a cascade-like model of the ribbon synapse 16 with different vesicle pool sizes, transfer rates and other synaptic properties. Exploiting recent 17 18 developments in simulation-based inference, we obtain full posterior estimates for the parameters and compare these across different retinal regions. The model enables us to 19 extrapolate to new stimuli and to systematically investigate different response behaviours of 20 21 various ribbon configurations. We also provide an interactive, easy-to-use version of this 22 model as an online tool. Overall, we show that already on the synaptic level of single neuron 23 types there exist highly specialized mechanisms which are advantageous for the encoding of different visual features. 24

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AUTHOR CONTRIBUTIONS. The manuscript was conceptualised and developed by CS, TY, TB and PB. CS established the models and together with JO adapted the inference method. TY performed electron microscopy and two-photon imaging, pre-processing of the data and generated transgenic lines, with support from TB. CS performed additional analyses and statistical testing, with help from PB. TB and PB supervised the project. The manuscript was written by CS and TB with input from TY and PB. Funding was acquired by TB, PB and TY.

#### 45 INTRODUCTION

Ribbon-type synapses feed high-bandwidth sensory signals into their 46 postsynaptic networks (reviewed in e.g. (Moser et al., 2019; Sterling and 47 48 Matthews, 2005)). However, depending on the species, modality, or receptor type the nature of this synaptic transfer can differ greatly. For example, 49 auditory systems typically operate at higher frequencies than visual systems. 50 and accordingly auditory inner hair cells tend to use "faster" ribbon synapses 51 compared to those of photoreceptors (Baden et al., 2013a; Moser et al., 52 53 2019). Moreover, amongst photoreceptors of the vertebrate eye, rods and 54 cones tend to differ greatly in the way they use their ribbons (Regus-Leidig 55 and Brandstätter, 2012; Sterling and Matthews, 2005). Rods generally have large ribbons that can dock many hundreds of vesicles at a time, 56 57 concentrated at a single release site, to support focussed, low-noise transmission (e.g. (Hays et al., 2021)). In contrast, cones usually use multiple 58 59 smaller ribbons, often positioned at different release sites in a single pedicle, to serve diverse postsynaptic circuits (e.g. (DeVries et al., 2006; Jackman et 60 al., 2009)). The ribbon synapses in electrosensory organs of elasmobranchs 61 62 take such ribbon tuning to the extreme (Bellono et al., 2018). For example, 63 sharks achieve high-amplitude pulsatile transmission required for predation by combining greatly elongated ribbons with an ion-channel composition that 64 supports broad spiking. In contrast, skates drive their smaller ribbons using 65 graded voltage signals to support low-amplitude, oscillatory transmission 66 suitable for intraspecific communication. This suggests that ribbons and their 67 associated molecular machinery are important structural tuning sites of 68 synaptic function in many sensory systems. 69

- However, the functional requirements of synaptic transmission do not only 70 differ across neuron types, but also within (Baden et al., 2013b; Franceschini 71 et al., 1981; Sinha et al., 2017; Szatko et al., 2019; Yoshimatsu et al., 2020b; 72 73 Zimmermann et al., 2018). For example, in vision, different parts of the eye 74 survey different parts of visual space, often with distinct distribution of light and visuo-ecological significance (reviewed in (Baden et al., 2020; Land and 75 76 Nilsson, 2012)). Correspondingly, we hypothesised that within-type functional tuning of a sensory receptor neuron should also utilise the vast 77 tuning potential of its ribbon. 78
- 79 We explored this idea in the model of larval zebrafish UV-cone photoreceptors, which exhibit profound structural, molecular and circuit 80 differences depending on their location in the eye (Yoshimatsu et al., 2020b). 81 First, UV-cone density varies across retinal regions and peak in the acute 82 zone (AZ), and to a lesser extent also nasally (Yoshimatsu et al., 2020b; 83 Zimmermann et al., 2018). Furthermore, UV-cones in the AZ combine an 84 enlarged outer segment with molecular tuning of the phototransduction 85 cascade, an elevated calcium baseline and strong feedback from horizontal 86 cells to boost detection brighter-than-background stimuli. This likely supports 87 visual prey capture of UV-bright water-borne micro-organisms such as 88 89 paramecia. In contrast, UV-cones in other parts of the eye preferentially 90 respond to darker-than-background stimuli, which may serve silhouette

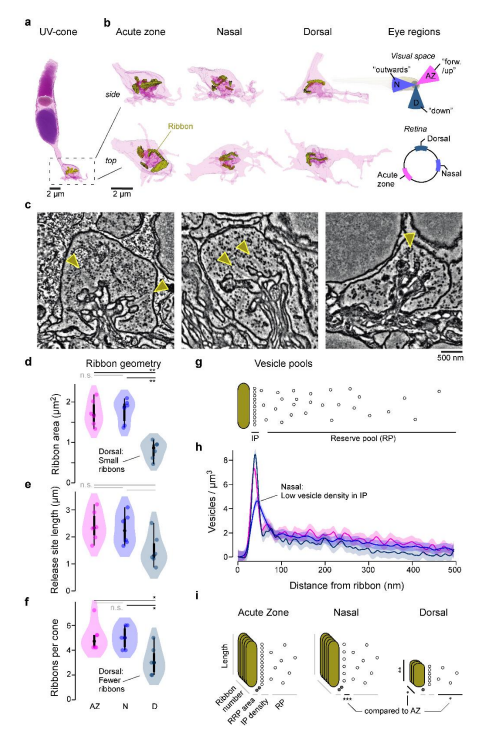
91detection of nearby objects against the backdrop of bright UV-scatter from92the sun. Amongst themselves, non-AZ UV cones further differ in additional93aspects, including their absolute light sensitivity. Building on these earlier94results, we here asked if and how the actual synaptic transfer differs amongst95UV-cones across the eye.

- First, we used electron microscopy to reveal eve-region specific structural 96 differences amongst UV-cone ribbons and their presynaptic distribution of 97 vesicles. Next, we found kinetic differences in synaptic transfer using in-vivo 98 99 simultaneous dual-colour 2-photon imaging of the same pedicles' 100 presynaptic calcium and resultant release. We then tied these findings 101 together in a biophysical model, which enables computationally exploring possible underlying biological mechanisms and sites of tuning within the 102 release cascade. Finally, we generalised our findings into an online model 103 of synaptic transfer from the ribbon that enables free control over all key 104 105 parameters, including ribbon dimensions, their dynamics, and the behaviour 106 underlying calcium drive (available of online at http://www.tinyurl.com/h3avl1ga). 107
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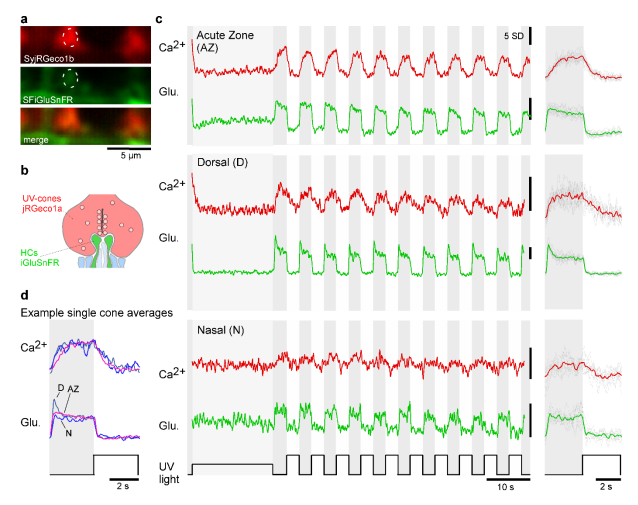
## RESULTS

UV-cone ribbon geometry and vesicle distributions differ with eye 110 position. To establish possible structural differences amongst larval 111 112 zebrafish UV-cone ribbon synapses, we obtained volumetric electron 113 microscopy datasets of the outer retina taken from three different regions: Acute zone (AZ), Nasal (N) and Dorsal (D). For each region, we anatomically 114 identified (Yoshimatsu et al., 2020a) and 3D-reconstructed UV-cone 115 pedicles ( $n_{AZ, N, D} = 6, 6, 6$ ) including their full complement of ribbons and 116 surrounding vesicles (Fig. 1a-c, Methods). This revealed that dorsal ribbons 117 were smaller (Fig. 1d,e) and less numerous (Fig. 1f) compared to AZ or nasal 118 ones. However, nasal UV-cones had the lowest vesicle density immediately 119 120 adjacent to the ribbon (Fig. 1g,h, Fig. S1). In addition, further away from the 121 ribbon, the vesicle density was lowest in dorsal UV-cones, and highest in AZ 122 UV-cones (Fig. 1h, Fig. S1). Although it is possible that the experimental procedure distorted the vesicle distribution slightly, it is unlikely that this 123 effect is disproportionately prominent in one eye region compared to the 124 others. Taken together, the overall complement between ribbon number, 125 geometry and vesicle distributions therefore markedly differed across the 126 three regions of the eye (Fig. 1i). We next asked if and how these structural 127 differences may translate into differences in synaptic function. 128





130 Figure 1. Eye-region-specific structural tuning of a ribbon synapse in UV-cones. a, Example of a full UV-cone 131 reconstruction, taken from the acute zone. Dark purple: nucleus, light-purple: mitochondria, purple: outer segment, 132 yellow: ribbon. b, Zoom-ins of UV-cone terminals from different regions, which are illustrated on the rightmost 133 panels. Ribbons are highlighted in yellow. Each terminal is shown from the side (top) and from below (bottom). c. 134 Example electron-microscopy images from each zone, with arrowheads indicating ribbons. d,e,f Violin plots of the 135 ribbon geometry and number for the three different regions (two-sided shuffling test with Bonferroni correction, n<sub>AZ</sub>, 136  $n_N$ ,  $n_D = 6$ , 6, 6, \* p<0.05, \*\* p< 0.01). **g**, Two-dimensional schema of the vesicle pools at a ribbon synapse. **h**, 137 Mean and 95% confidence intervals for vesicle densities as a function of distance to the ribbon. Predictions are 138 made from a Generalized Additive Model (GAM, Methods), (see Fig. S1 for statistical comparisons). i, Summary 139 schema of observed EM-level differences between UV-cones at the level of ribbon geometry, number, and vesicle 140 distributions. Asterisks indicate significant differences compared to AZ. The stacked ribbons (gold) indicate the 141 ribbon number per cone, whereas the vesicles (small circles) are exemplified in a two-dimensional plane for a single 142 ribbon.

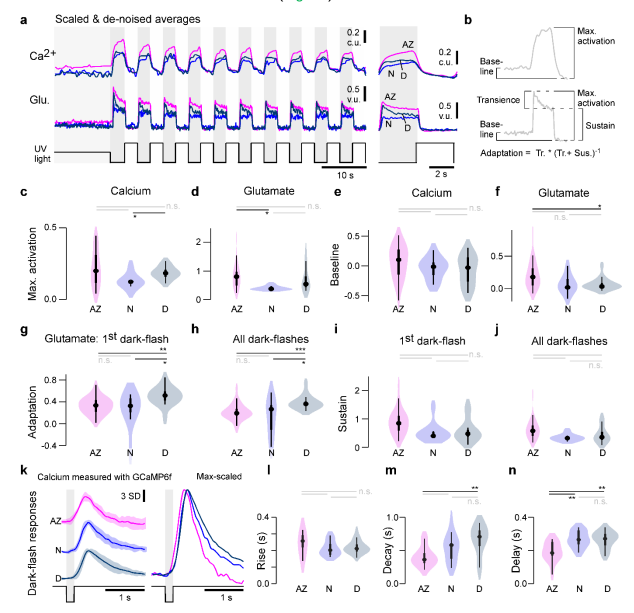


144 Figure 2. Simultaneous in vivo imaging of synaptic calcium and release. a,b Simultaneously acquired two-145 photon scans of cone terminals and opposing horizontal cell dendrites, with cone-pedicles expressing SyjRGeco1b 146 (red), and horizontal cell dendrites expressing SFiGluSnFR (green), and schematic representation, showing the 147 cone-pedicle (red) with ribbon and vesicles, as well as horizontal cell processes (green) and bipolar cell dendrites 148 (blue). c, Examples of raw calcium (red) and glutamate (green) traces recorded simultaneously from single UV-149 cones, one from each eye region as indicated. The averaged traces, and superimposed stimulus repetitions are 150 shown on the right d, Overlay of the averaged traces in (c), highlighting different glutamate responses despite very 151 similar calcium responses.

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152 Eve-region dependent differences in UV-cone release kinetics. To 153 simultaneously monitor presynaptic calcium and resultant glutamate release from single UV-cone pedicles in vivo, we expressed the "green" fluorescent 154 glutamate biosensor SFiGluSnFR in horizontal cells postsynaptic to the 155 cones, and the "red" calcium biosensor SyjRGeco1b in all-cones (Fig. 2a,b, 156 Methods). Biosensor expression appeared to be uniform across the eye, in 157 line with previous work (Yoshimatsu et al., 2020b). Furthermore, there was 158 no obvious spectral mixing of the two fluorescence channels (SFig. 2a,b). 159 160 UV-cones were unambiguously identified based on their robust responses only to UV-light (Methods, (Yoshimatsu et al., 2020a)). We then concurrently 161 162 recorded red and green fluorescence signals under two-photon during presentation of 100% contrast widefield flashes of UV-light (3 s On, 3 s Off), 163 starting from a 50% contrast background (Fig. 2c, Methods). In example 164 recordings from each eye region, this revealed very different glutamate 165 signals during light offsets, despite similar appearing calcium signals (Fig. 166

2c). When scaled to their common sustained component, the dorsal cone was much more transient compared to the nasal cone, with a kinetically intermediate AZ-cone (Fig. 2d).



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Figure 3. Physiological differences in light responses between UV-cones from different eye-regions. a, 171 172 Scaled and de-noised calcium and glutamate recordings averaged across multiple ROIs (Methods). We refer to the 173 scaling as calcium units (c.u.) and vesicle units (v.u.), as the same traces also serve as input for the biophysical 174 model (see Fig. 4). b, Schema of the calculated indices in (c-j) (Methods). The transience index is computed as max.– sustain . c-j, Quantification of physiological differences for the three different retinal regions (two-sided shuffling 175 max. test with Bonferroni correction, n<sub>AZ</sub>, n<sub>N</sub>, n<sub>D</sub> = 30,9,16, \* p<0.05, \*\* p< 0.01, \*\*\* p<0.001). k, GCaMP6f recordings 176 177 from (Yoshimatsu et al., 2020b). Mean±SD and overlaid mean traces in response to a 200 ms dark flash stimulus. 178 I-n, Quantification of physiological differences for the GCaMP6f recordings: time constants for an exponential rise, 179 decay as well as delay time to response (see also methods) (two-sided shuffling test with Bonferroni correction, 180  $n_{AZ}$ ,  $n_N$ ,  $n_D = 13,17,22$ , \* p < 0.05, \*\* p < 0.01, \*\*\* p < 0.001).

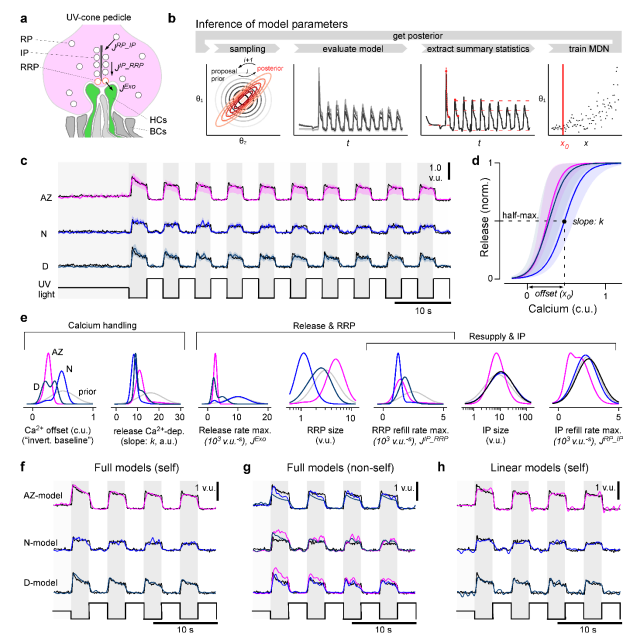
181To systematically test for consistent differences between synaptic transfer in182the three eye regions, we recorded paired calcium and glutamate signals183from a total of n = 30, 16, 9 AZ, dorsal and nasal UV-cones from n = 3, 4, 4184fish, respectively (Fig. S2c,d). We then scaled and de-noised each recording

(Methods) and computed the mean traces (Fig. 3a) as well as key 185 parameters relating to the amplitudes and kinetics of calcium and glutamate 186 signals (Fig. 3b-j). For this, all traces where scaled such that the UV-bright 187 stimulus intervals had a zero mean and standard deviation of one. We 188 previously showed that these intervals correspond to the lowest possible 189 calcium and glutamate release at the stimulus brightness used (Yoshimatsu 190 191 et al., 2020b) and can therefore be used as a common baseline across 192 zones. We present the rescaled traces (Fig. 3a) in c.u. (calcium units) and 193 v.u. (vesicle units) respectively, to be consistent with the used units in the 194 model later. Based on the rescaled traces we computed several indices of calcium and glutamate response amplitudes and their kinetics (Fig. 3b), 195 which together confirmed and extended initial observations from the single-196 197 pedicle examples (cf. Fig. 2). In particular, both at the level of calcium (Fig. 3c) and glutamate (Fig. 3d), nasal UV-cones exhibited small peak-198 amplitudes. In addition, the glutamate release of AZ UV-cones was 199 increased during the 50% contrast period at the start of the stimulus (Fig. 3f). 200 201 In line with previous work, the AZ calcium baseline also appeared elevated 202 (Fig. 3a), however this difference was not statistically significant (Fig. 3e). This was likely related to the lower signal-to-noise ratio of jRGeco1b signals, 203 compared to those of GCaMP6f as used previously (Yoshimatsu et al., 204 2020b). Next, we quantified amplitudes of transient and sustained 205 components at the level of glutamate (Fig. 3b,g-i). For this, we analysed the 206 207 first flash response separately from the mean of subsequent ones because all regions exhibited notably stronger adaptation during the first flash (Fig. 208 209 3a). Overall, this consistently revealed the most pronounced within-pulse 210 adaptation in dorsal UV-cones (Fig. 3g,h), but no significant differences in 211 the sustained components (Fig. 3i,j).

To investigate the differences in presynaptic calcium in more detail, we 212 reanalysed previously published data of calcium recordings with 213 SyGCaMP6f in response to a 200 ms "dark flash" (Yoshimatsu et al., 2020b). 214 The kinetics of calcium responses were similar to each other across the three 215 zones (Fig. 3k), thus broadly supporting our previous results based on 216 jRGeco measurements. First, we found similar rise kinetics across all zones 217 (Fig. 3I) which therefore unlikely linked to the differences observed at the 218 219 level of glutamate. Nevertheless, the response in the AZ was weakly but significantly advanced (i.e. it occurred earlier) compared to nasally or 220 221 dorsally (Fig. 3n). Moreover, decay kinetics were significantly faster in AZ cones compared to dorsal cones (Fig. 3m) contrary to the adaptation index 222 of the glutamate recordings (Fig. 3g, i), hinting that the release dynamics 223 were shaped differentially by the synaptic machinery across zones. 224

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<sup>225</sup>Taken together, our results so far highlight a range of structural (Fig. 1) and226functional (Figs. 2,3) differences in the synaptic machinery of UV-cones227across different regions of the eye.



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231 Figure 4. A model of calcium evoked release from the ribbon. a, Schema of the movement of vesicles at a 232 ribbon synapse. The vesicles move from the reserve pool (RP) to the intermediate pool (IP) at the ribbon and finally 233 to the readily releasable pool (RRP) close to the membrane before they are released into the synaptic cleft to 234 activate the dendrites of invaginating horizontal cells (HC). b, Schema of the parameter inference: Over several 235 rounds, samples are first drawn from a prior, the model is then evaluated and summary statistics on which the 236 relevant loss function is calculated are extracted. Based on these values and the sampled parameters a mixture of 237 density network (MDN) is trained and evaluated to get the posterior / prior for the next round. c, Model predictions 238 for the three regions (mode and 90% prediction intervals from posterior samples in colour, data in black). d, Mode 239 and 90% prediction intervals of the model's non-linearity. e, Prior and one-dimensional marginals of the posterior 240 distributions (see Fig. S4a for the two-dimensional marginals). The units are vesicle units (v.u.) and calcium units 241 (c.u.), referring to the scale invariance of the model. f, Evaluation of the best region-specific models on its region-242 specific calcium traces. g, Evaluation of best region- specific model on the calcium races of the other regions. h, 243 Evaluation of the linear baseline model (model in colour, data in black): especially the transient components are 244 missed. See Fig. S3b for a quantification of the goodness of the fits.

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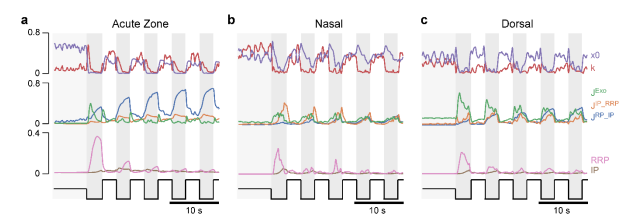
A model of glutamate release at the ribbon synapse. To systematically explore the possible mechanistic basis of the observed eye-wide variations in UV-cone synaptic functions, we next modelled the release machinery of 248 the ribbon with a biophysical interpretable model that converts calcium signals to glutamate release (Baden et al., 2014; Schröder et al., 2019). The 249 250 model consisted of three different vesicle pools (reserve pool (RP), intermediate pool (IP) and readily releasable pool (RRP)), the changing rates 251 between these pools ( $J^{RP_{-}IP}$ ,  $J^{IP_{-}RRP}$ ,  $J^{Exo}$ ) and a sigmoidal non-linearity with 252 slope k and offset  $x_0$  which converts the calcium concentration into the final 253 254 glutamate release (Fig. 4a, Methods). Building on recent advances in 255 simulation-based Bayesian inference (Gonçalves et al., 2020; Lueckmann et 256 al., 2017), we estimated posterior distributions over the model parameters 257 for each of the region-specific datasets. In summary, the inference methods iterated the following steps over several rounds (Fig. 4b): First we draw 258 samples from a prior distribution, evaluated the model and extracted 259 summary statistics on which the relevant loss was computed. Based on 260 these loss values and the sampled parameters a mixture of density network 261 was trained and finally evaluated to get the posterior of one round which was 262 used as the prior for the next round. For the summary statistics, we defined 263 264 features which captured all essential components of the release dynamics, 265 such as transient and sustained components or peak heights (Fig. 4b, Methods). The model fitted the functional data well and accurately modelled 266 the pronounced first UV-flash response differences between zones (Fig. 4c). 267 From here, the inferred posteriors (see Fig. S4a for two dimensional 268 marginals) allowed us to compare the likely parameters between the different 269 zones, including the estimated uncertainties. This allowed pinpointing 270 possible key differences: For example, the calcium offset ( $x_0$ , which can be 271 understood as the inverted calcium baseline, Methods) was markedly 272 273 increased in the nasal model (Fig. 4d,e). In contrast, in case of maximal 274 release rate, the posterior for the nasal data stayed close to the prior, indicating that this parameter was not essential to reproduce the traces. For 275 RRP sizes and associated maximal refill rates the model required the 276 277 smallest values nasally (Fig. 4e). The posteriors for IP size and associated refill rates were rather broad and did not allow to identify regional differences. 278 279 Interestingly, with few exceptions (e.g. the maximal release rate and calcium offset), the parameter posteriors were mostly uncorrelated (Fig. S4a), 280 indicating that there is only little structure in the optimal parameter 281 282 landscape. This also suggests that the model and data leave little room for possible compensatory mechanisms, as this would result in a clear 283 correlation of the involved parameters. 284

To confirm that the different model outputs do not simply rely on differences 285 in the calcium inputs but rather on the differences of the inferred parameters, 286 we compared the performance of the zone-specific models by shuffling the 287 inferred parameters and glutamate datasets pairwise across zones (Fig. 4f,g, 288 Fig. S4b). The match between the model output and measured glutamate 289 290 release for a calcium input was generally worse for parameters corresponding to a different region, confirming that our release models were 291 indeed regionally specific. For example, the sustained component of the AZ 292 293 could not be captured by either the dorsal or nasal models (Fig. 4g). Moreover, the models produced different transient behaviours independent 294

295 of the calcium inputs (Fig. 4f,g). A quantification of this model comparison shows that the shuffled models achieved lower loss values when evaluated 296 on self-, compared to non-self calcium inputs (Fig. S4b,c). The model 297 differences were especially high on the relevant loss (Fig. S4b), which was 298 based on the summary statistics and paid special attention to features like 299 transiency. However, already the mean squared error (MSE) as loss function 300 301 confirmed this difference: the region-specific models evaluated on self 302 calcium inputs outperformed the models on non-self inputs (Fig. S4c). 303 Additionally, we compared the biophysical model to a statistical linear 304 baseline model (Methods). While as expected the linear model captured the general shape of the flash responses, it was not able to model glutamate 305 transients nor adaptation over several flashes (Fig. 4h). This also resulted in 306 much higher loss values for the relevant loss function compared to the best 307 308 biophysical model, which indicates the mismatch for essential features such 309 as transiency (Fig. S4b). The biophysical model also outperformed the linear model in terms of the MSE (Fig. S4c). Together, this indicates that our 310 311 modelling approach was sufficiently detailed for the posed problem, 312 suggesting that the posterior distributions of each regional model can usefully inform about the differences that underpin region-specific transfer 313 functions from synaptic calcium to release via the ribbon. 314

Predicting region-specific processes. An important strength of our 315 modelling approach was that it allowed systematically exploring the possible 316 influence of parameters such as vesicle pool sizes, their vesicle movements, 317 and their calcium dependence, on the model output. To this end, we 318 conducted a sensitivity analysis by computing the first order Sobol indices 319 (Methods), a measure of the direct effect of each parameter on the variance 320 of the model output. More specifically, it denotes the expected reduction in 321 relative variance of the model output if we fix one parameter. For the 322 computation of the Sobol indices, broadly speaking, a large number of 323 parameters were drawn from the posterior distribution and the model was 324 evaluated on these parameters. Afterwards the reduction in variance of the 325 model evaluations was computed if one dimension of the parameter space 326 was fixed (for details see Methods). The Sobol indices revealed a generally 327 high influence of the calcium parameters ( $x_0$  and k) during UV-bright periods 328 329 and around light-dark transitions in all three zones (Fig. 5a-c, top row). In contrast, beyond an initial key role of the RRP size for shaping the first dark-330 flash response, pool sizes generally played only relatively minor roles (Fig. 331 5, bottom row). Instead, the most obvious region-wise differences occurred 332 amongst vesicle transition rates between the pools. For example, the refilling 333 rate of the IP from the RP  $(J^{RP_{-}IP})$  was increasingly critical for shaping dark 334 flash responses in the AZ-model (Fig. 5a, blue) but had comparatively little 335 influence in the nasal model (Fig. 5b, blue). In contrast, the maximal release 336 rate  $(J^{Exo})$  particularly influenced the variance in early dark flash responses 337 in the dorsal model (Fig. 5c, green), while nasally refilling of the RRP from 338 the IP  $(J^{P_RRP})$  played a greater role (Fig. 5b, yellow). Together, this analysis 339 340 suggests that particularly the rates of vesicle transfer between pools, rather 341 than the pool sizes themselves or their calcium dependence, may underpin

342 343 the experimentally observed region wise differences in release properties from zebrafish UV-cones *in vivo*.



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Figure 5. Sobol Indices. a-c, Sensitivity of the model for the different model parameters measured by the Sobol
 Index (Methods). The Sobol Index measures the expected reduction in relative variance for the fixation of parameter
 θ<sub>i</sub>. It depends on the posterior distribution and is therefore different for the fits to the three regions.

General rules of ribbon tuning. We next sought to explore the general 348 349 parameter landscape underlying release from the ribbon. For this, we 350 calculated the same indices as in the corresponding data from 2P imaging (Fig. 3), but this time on the model output obtained by simulating the model 351 with parameters drawn from the posterior distributions. As expected, this 352 reproduced the trends previously measured in vivo, including the low 353 maximal activation nasally (Fig. 6a), the largest transient component dorsally 354 (Fig. 6b), and the largest sustained component in the AZ (Fig. 6c). 355

- From here, we simplified the model by fixing the slope of the calcium non-356 linearity (k) and defining vesicle change rates as fractions of the 357 corresponding pool sizes (Methods). To be able to stimulate the model with 358 359 arbitrary "light" stimuli, we moreover implemented a linear calcium model 360 based on a convolution with a biphasic kernel to reflect cone-activation by light (Schnapf and Baylor, 1987), and monophasic kernel to reflect calcium 361 kinetics (Baden et al., 2014). This latter kernel was varied in subsequent 362 simulations to explore the impact of calcium kinetics on synaptic 363 performance (Methods). Together, this allowed us to reduce the parameter 364 space while at the same time identifying underlying computational principles. 365 In the following, we always included the three fitted eve-region specific 366 parameter sets as a point of comparison (coloured "dots" on top of heatmaps 367 in Figs. 6,7). These dots should be treated with some caution, since in the 368 simplified model they do not necessarily match the original ones in every 369 dimension. 370
- Exploring this model (Fig. 6d-f), we found that both maximal activation (Fig. 6d) and the size of the sustained component (Fig. 6f) could be tuned by varying RRP and/or IP pool sizes (left column), with negligible contributions from the maximal release rate or the calcium offset (right column). In contrast, the transient component primarily hinged on the maximal release rate (Fig. 6e, right column), with more complex additional contributions from the interplay of vesicle pool sizes (left column). Accordingly, our generalised

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model suggests that transient and sustained responses can be defined largely non-overlapping properties of the ribbon, possibly providing a powerful handle in for their independent tuning. For further exploration, the full model is available online as an interactive tool (Fig. 6g) via *google colab* (<u>http://www.tinyurl.com/h3avl1ga</u>) or on *github* (<u>https://github.com/coschroeder/cone\_ribbon</u>).

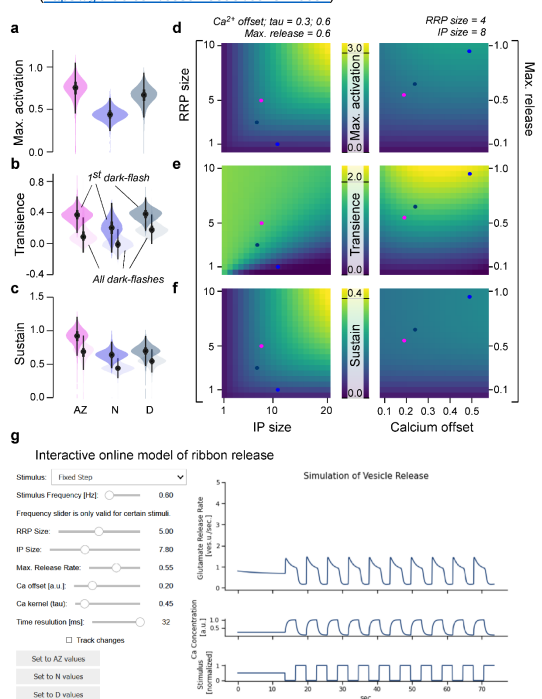
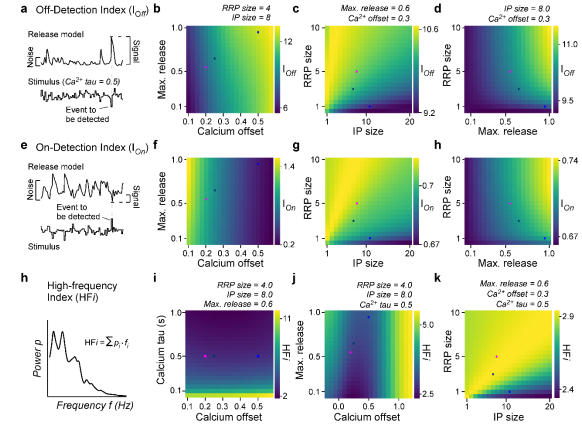




Figure 6. General rules of ribbon tuning: Basic response parameters. a-c. The same indices as in Fig. 3, but here calculated on 10,000 model evaluations on samples from the posteriors. The model has learned the differences between the retinal regions and reproduces these differences (see Fig. 3 for comparison). d-f, Indices as in (a-c), calculated on different parameter combinations as indicated. For this analysis, a step-stimulus feeding into a linear calcium model was added as the input to the release model (Methods). For definition of the indices see also Fig. 3b. g, Screenshot of the interactive online model, available at http://www.tinyurl.com/h3avl1ga.

Frequency dependence and event detection. In a final step, we explored the model behaviour on new stimuli and investigated the influence of the model parameters on different coding properties of the synapse. First, we measured the detectability of a high amplitude dark-event amongst an otherwise noisy stimulus sequence (Fig. 7a, Methods). This highlighted the calcium offset ( $x_0$ ) as a key parameter (Fig. 7b). Once  $x_0$  is set, additional benefits could be gained from increasing RRP-size but only small benefits from increasing IP size (Fig. 7c), ideally in further combination with a high maximal release rate (Fig. 7d). We next measured the detectability of lightevents in the same way (Fig. 7e, Methods). This showed that beyond an inverse dependence on calcium baseline (Fig. 7f cf. Fig. 7b), the ribbon parameters that benefitted the detection of On- and Off-events were in fact virtually identical (Fig. 7g,h, cf. Fig. 7c,d). The low calcium offset (i.e. high baseline) supporting the detection of On-events is in line with our previous work (Yoshimatsu et al., 2020b), where the AZ showed highest calcium baseline and an enhanced ability to detect visuo-ecologically important UV-On events such as the presence of prey.



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Figure 7. General rules of ribbon tuning: Event detection and high-frequency encoding. a, An Off-Detection index (*l*<sub>off</sub>) measures the model's baseline-noise normalised response amplitude to an Off-event in the stimulus as indicated (Methods). b-d, *l*<sub>off</sub> for different parameter combinations. e-h, as (a-d), but for an On-detection index (*l*<sub>on</sub>).
h, The high-frequency index (HFi) is a weighted sum of the discretized power spectrum and indicates the behaviour in a high frequency regime. i-k, HFi for different parameter combinations. The fixed parameters are shown as titles in each panel. Note the different colour scales between panels. Further note that in (j) we explored the x<sub>0</sub> parameter space up to extreme response behaviour, which may not be physiologically plausible.

416 Finally, we explored how well different ribbon models could transmit fast 417 temporal flicker, here summarised by a high-frequency index (HF*i*) (Fig. 7h, 418 Methods). This revealed that this property primarily depended on the calcium 419 kinetics, rather than the specific tuning of the ribbon itself (Fig. 7i-k). Beyond 420 calcium dynamics (here, fixing the time constant *r* for calcium at 0.5 s), again 421 the calcium offset (Fig. 7j) as well as an approximate balance of medium-422 sized pools for both the RRP and IP (Fig. 7k), could provide additional 423 support for encoding high-frequency components.

### 424 DISCUSSION

- Combining ultrastructural evidence (Fig. 1), in vivo dual-colour 2-photon 425 imaging (Figs. 2,3) and computational modelling (Figs. 4,5) we have shown 426 427 how ribbon synapses belonging to the same neuron type can be regionally 428 tuned to support distinct synaptic transfer functions depending on their 429 location in the eye. Our findings complement and extend our recent demonstration that also upstream properties of these UV-cones, are 430 regionally tuned to support different visuo-ecological functions (Yoshimatsu 431 et al., 2020b). We then further generalised this model to explore how specific 432 properties ribbon function can be principally used and traded off against one 433 another to achieve a broad range of synaptic properties (Fig. 6,7). 434
- Linking ribbon structure and function. Our findings that both UV-cone 435 ribbon ultrastructure (Fig. 1) and their effective synaptic transfer (Fig. 2,3) 436 437 systematically differed between regions supports the notion that these two 438 sets of properties are linked (Regus-Leidig and Brandstätter, 2012; Sterling 439 and Matthews, 2005; Wichmann and Moser, 2015). However, building a 440 direct bridge between them remains difficult. This difficulty is part related to the absence of a direct experimental link between our EM- and 2P-datasets, 441 which we did not attempt in view of its extreme technical challenges (Holler 442 443 et al., 2021) that to our knowledge have not been overcome for any ribbon synapse. Moreover, there is often more than one possible interpretation 444 between a finding from EM and its functional consequence. For example, the 445 446 size of a ribbon is expected to be linked to the number of vesicles it can hold. 447 However, this link presumes that ribbon-attached vesicle density is fixed, 448 which may not be the case, for example due to conceivable variations in available binding sites, or amongst vesicle-transfer rates between pools (e.g. 449 Fig. 4). 450
- Such possible complexity is illustrated in case of UV-cone regional 451 variations: For example, dorsal ribbons were by far the smallest and least 452 numerous (Fig. 1d.f). From here, it is tempting to speculate that therefore 453 their effective IP size, and perhaps also their RRP size, might be 454 comparatively small. However, dorsal UV-cones also had a particularly high 455 456 vesicle density near the ribbon, while instead nasal UV-cones had markedly reduced vesicle density here despite their otherwise large ribbons (Fig. 1g, 457 Fig. S1). This strongly suggest that looking at ribbon geometry alone may be 458 insufficient to accurately predict effective pool sizes. Instead, joint 459 consideration of ribbon area and vesicle density may be more informative as 460 supported by our model, which produced a near-identical IP size and 461 generally low RRP sizes for both nasal- and dorsal UV-cones (Fig. 4e). In 462

463 agreement, AZ-cones combined large ribbons with a high vesicle density near the ribbon, and in their case the model did predict the largest RRP size 464 (though a similar IP size). Although the differences of RRP sizes which were 465 inferred by the model did not match the anatomical numbers quantitatively, 466 we were able to infer the trends of the differences in the anatomical 467 structures from pure functional data. Notably, model IP sizes generally 468 stayed near the prior (Fig. 4e) and exhibited low Sobol indices (Fig. 5), which 469 470 suggests that this property was not critical for explaining the glutamate 471 responses to our relatively slow and simple test stimulus which was necessitated by the generally low SNR of jRGeco1b signals (Fig. 2). It 472 remains possible that regionally distinct IP-size estimates would emerge in 473 474 response when fitted to responses to different stimuli.

- Beyond ribbon sizes, our ultrastructural analysis further highlighted an 475 elevated vesicle density further away from the ribbon in AZ UV-cones (Fig. 476 477 1g, Fig. S1), potentially indicative of an increased RP and/or an increased 478 rate of IP refilling. However, the model produced only minor differences in IP refill rates between regions, but in the reversed order (Fig. 4e), and for 479 480 simplicity RP size was fixed because it did not strongly affect function within 481 the tens of seconds timescales interrogated (Baden et al., 2014). Accordingly, a direct link between UV-cone RP vesicle density and function 482 remains outstanding. Conceivably, this property could be helpful for 483 supporting the particularly high total vesicle turnover in AZ-cones (Fig. 3d,f,i) 484 at longer timescales. 485
- At the level of measured function, dorsal cones stood out in that their release 486 487 was particularly transient (Fig. 3a, g, h) as would be suited for pulsatile, rather than continuous transmission. Their generally small but densely populated 488 ribbons and small RP may be well suited to support this property. In possible 489 490 agreement, the model predicted a faster RRP refill rate in dorsal cones, 491 which might be useful to ensure the RRP is rapidly replenished after each 492 pulse. Conversely, the effective absence of any transient component nasal UV-cones (Fig. 3a,q,h) in the model resulted in the lowest RRP refill rate, 493 494 which may appear counterintuitive, but paired with a small nasal RRP resulted in the sustained response behaviour observed in Fig. 4f. 495 Additionally, this may further link to the abovementioned low IP occupancy 496 in nasal UV-cones, as observed under EM (Fig. 1g). Moreover, the nasal 497 model simultaneously predicted the smallest RRP but the largest RRP 498 release rate, which resulted in a situation where the very few vesicles ready 499 for release were immediately dumped, thus preventing the build-up of 500 vesicles that enable the transients of dorsal and AZ UV-cones. Importantly, 501 502 most functional differences across retinal regions appeared at the level of 503 glutamate, whereas only subtle differences in the dynamics were observed at the level of calcium. The faster decay in AZ (Fig. 3m) is linked to horizontal 504 cells as reported previously (Yoshimatsu et al., 2020b), and the temporally 505 506 advanced calcium signals measured in the AZ by GCaMP6f (Fig. 3n) cannot 507 explain the differences in the glutamate dynamics. Moreover, while it remains possible that the iGluSnFR recordings slightly distorted the time-508

509 courses of glutamate signals, it is unlikely to strongly affect our conclusions, 510 because any such effects would apply equally to all measured eye regions.

- 511Taken together, while it remains difficult to make definitive links between512ribbon structure and function, a number of inferences can be drawn which513may usefully inform our understanding of the ribbon's role in moulding514synaptic transmission to specific needs.
- 515 In the future it will be interesting to explore to what extent the differential ribbon tunings described here for larvae are also a characteristic of UV-516 cones in adult zebrafish. Adults feature a crystalline cone mosaic (reviewed 517 518 in (Baden, 2021)), meaning that numerical anisotropies in cones as they 519 occur in larvae (Zimmermann et al., 2018) are not expected. However, this 520 does not preclude the possibility that UV-cones have different properties across the retina. While adult zebrafish display a much broader array of 521 visual behaviours compared to larvae, their visual ecology remains poorly 522 explored, which makes it difficult to predict what UV-cone tunings - if any -523 524 might be expected.
- Model and simulation-based inference. The presented model (Fig. 4a) is 525 a modified version of the basic framework used in (Baden et al., 2014). Here 526 we extended the approach by combining simulation-based inference with 527 528 simultaneous calcium and glutamate recordings to extensively expand the 529 analysis of the model. Moreover, rather than modelling discrete vesicle 530 movements as in (Schröder et al., 2019), our continuous model enabled the use of available software toolboxes (Methods). With the simulation-based 531 532 inference approach (Fig. 4b) we obtained full posterior distributions (Fig. 4e) rather than point estimates which can lead to overconfident or incorrect 533 conclusions in the case of an under-constrained model or "sloppy" model 534 parameters. At the same time, posterior estimates can be seen as a global 535 method to identify sloppy and stiff parameters (in contrast to local methods 536 537 as in (Gutenkunst et al., 2007)), and additionally allowed us to conduct a 538 sensitivity analysis by computing the first Sobol indices (Fig. 5). These focus 539 on the variance of the model output rather than on the variance of the distributions and showed zone dependent, 540 posterior biologically interpretable time courses. This highlighted a time varying dependence of 541 the release on the model parameters. It indicates that already a simple step 542 stimulus is sufficient to show the influence of different anatomical properties 543 on features of the signal. Finally, by establishing an interactive online model 544 we encourage further exploration of the model and testing of hypotheses. 545
- 546General rules of ribbon tuning. Bringing together the model and observed547differences in response behaviour, we tested our model on new stimuli and548computed indices (Fig. 7) which might be relevant to a broad range of549sensory scenarios. For all investigated properties, we observed that the550baseline and dynamics of the calcium signal were critical (Fig. 7b,f,i),551whereupon the anatomical ribbon properties allowed fine-tuning the final552output behaviour (Fig. 7c,g,k).

553 Interestingly, the influence of the calcium baseline for the detection of highfrequency events followed a bimodal distribution (Fig. 7j) with intermediate 554 values offering the poorest performance. In combination with the 555 approximately equal and opposite effects of calcium baseline on the 556 detectability of On- and Off-events (Fig. 7b,f), this suggest that the calcium 557 baseline may present a key variable that enables ribbons to trade-off the 558 transmission of high frequency stimuli against providing an approximately 559 560 balanced On- and Off- response behaviour. Vice versa, it also suggests that 561 the transmission of high-frequency events benefits from the use of a highly 562 non-linear synapse that is either balanced to On- or Off-events, but not both. Finally, and perhaps unsurprisingly, the time-course of calcium decay was 563 pivotal for defining the possible working range of high-frequency 564 transmission regardless of the properties of the ribbon itself (Fig. 7i, cf. Fig. 565 7j,k). This effect was especially strong for a fast calcium decay (< 100 ms) 566 which is generally associated with nanodomains (Jarsky et al., 2010) rather 567 than decay dynamics >>100 ms for microdomains (Beaumont et al., 2005). 568 569 To what extent nano- or microdomain signalling dominates in larval zebrafish 570 UV-cones remains untested. However, in view of their similarity in pedicle architecture to mammalian rods (e.g. small, single invagination site), it 571 seems likely that also here already low micromolar calcium concentrations 572 that are typically associated with microdomains can evoke substantial 573 release (Thoreson et al., 2004). In any case, a primary determinant for tuning 574 575 a ribbon's high-frequency response is the local stimulus-driven calcium environment around the ribbon, rather than properties of the ribbon itself (see 576 577 also (Baden et al., 2014, 2013a)). Nevertheless, once this is set, large pool 578 sizes paired with a high release rate were generally preferable for all explored forms of signal detection. Taken together, our model therefore 579 suggests that while the pre-synaptic calcium is a critical variable, even with 580 a fixed calcium model the ribbon can be shifted into different response 581 behaviours. 582

- 583The effect and importance of calcium handling has also been shown in584ribbon synapses of inner hair cells, where even within an individual synaptic585compartment a local variation in calcium channels can lead to different586transfer functions at different release sides (Özçete and Moser, 2021).587Similar to the presented differences in UV-cones, such local heterogeneity588in inner hair cells might help to diversify the sensory signal and highlight589different features for downstream neurons.
- 590 591

### 592 METHODS

593 Animals. All procedures were performed in accordance with the UK Animals (Scientific Procedures) act 1986 and approved by the animal welfare 594 committee of the University of Sussex. Animals were housed under a 595 standard 14:10 day/night rhythm and fed three times a day. Animals were 596 grown in 0.1 mM 1-phenyl-2-thiourea (Sigma, P7629) from 1 dpf to prevent 597 melanogenesis. For 2-photon in-vivo imaging, zebrafish larvae were 598 immobilised in 2% low melting point agarose (Fisher Scientific, BP1360-599 600 100), placed on a glass coverslip and submerged in fish water. Eye 601 movements were prevented by injection of a-bungarotoxin (1 nL of 2 mg/ml; 602 Tocris, Cat: 2133) into the ocular muscles behind the eye.

- 603For all experiments, we used 6-7 days post fertilization (dpf) zebrafish (Danio604rerio) larvae. The following previously published transgenic lines were used:605Tg(cx55.5:nlsTrpR), and Tg(tUAS:SFiGluSnFR) (Yoshimatsu et al., 2020b).606In addition, Tg(gnat2:SyjRGco1a) lines were generated by injecting607pTol2CG2-gnat2-SyjRGeco1a, plasmids into single-cell stage eggs. Injected608fish were out-crossed with wild-type fish to screen for founders. Positive609progenies were raised to establish transgenic lines.
- 610 The plasmid was made using the Gateway system (ThermoFisher, 611 12538120) with combinations of entry and destination plasmids as follows: 612 pDESTtol2CG2 (Kwan et al., 2007), p5E-gnat2 (Lewis et al., 2010; Yoshimatsu et al., 2016), pME-SyjRGeco1a, p3E-pA. Plasmid pME-613 SyjRGeco1a was generated by inserting a polymerase chain reaction (PCR)-614 amplified jRGeco1a (Chen et al., 2013; Dana et al., 2016) into pME plasmid 615 and subsequently inserting a PCR amplified zebrafish synaptophysin without 616 stop codon at the 5' end of jRGeco1a. 617
- Electron Microscopy. We used a previously published EM dataset of the 618 larval zebrafish outer retina for this study (Yoshimatsu et al., 2020a). In the 619 original paper, we only used one image stack from the acute zone, but here 620 621 we have in addition included two further stacks from nasal and dorsal 622 regions, respectively. Image stacks were concatenated and aligned using 623 TrackEM (NIH). The cones and ribbons were traced or painted using the tracing and painting tools in TrackEM2 (Cardona et al., 2012). Vesicle 624 density on the ribbons was measured in 6 representative sections per cone. 625 We selected sections where ribbons were aligned perpendicular to the 626 sections. Ribbon release site length and area were measured in the 3D 627 reconstruction of the ribbons. 628
- **2p imaging.** All 2-photon imaging was performed on a MOM-type 2-photon 629 630 microscope (designed by W. Denk, MPI, Martinsried; purchased through 631 Sutter Instruments/Science Products) equipped with a mode-locked Ti:Sapphire laser (Chameleon Vision-S, Coherent) tuned to 980 nm. We 632 633 used two fluorescence detection channels for iGluRSnFR (F48x573, AHF/Chroma) and jRGeco1a (F39x628, AHF/Chroma), and a water 634 immersion objective (W Plan-Apochromat 20x/1,0 DIC M27, Zeiss). For 635 image acquisition, we used custom-written software (ScanM, by M. Mueller, 636

MPI, Martinsried and T. Euler, CIN, Tuebingen) running under IGOR pro 6.3
for Windows (Wavemetrics). Recording configuration was 124 x 32 pixels
(2ms per line, 15.6 Hz). Light stimuli were delivered through the objective,
by band-pass filtered light emitting diodes (LEDs, 'red' 588 nm, B5B-434-TY,
13.5cd, 8°, 20 mA; 'green' 477 nm, RLS-5B475-S; 3-4 cd, 15°, 20mA; 'blue'
415 nm, VL415-5-15; 10-16 mW, 15°, 20 mA; 'ultraviolet, UV' 365 nm,
LED365-06Z; 5.5 mW, 4°, 20 mA, Roithner, Germany).

- 644 All LEDs were jointly further filtered using FF01-370/36 (AHF/Chroma) and 645 synchronized with the scan retrace at 500 Hz using a microcontroller as 646 described in (Zimmermann et al., 2020). The LED intensity was 1.10<sup>5</sup>, and 647 2.10<sup>5</sup> photons per cone per second for adaptation period and UV flash, respectively, which corresponds to a low-photopic regime. A stimulus time 648 649 marker embedded in the recording data was aligned the traces with a temporal precision of 2 ms. For all experiments, animals were kept at 650 constant background illumination for at least 5 s at the beginning of each 651 recording to allow for adaptation to the laser. Regions of interest (ROIs), 652 corresponding to individual presynaptic terminals of UV-cones were defined 653 654 manually. For calcium, we restricted ROIs within 1 µm from the release site 655 at a terminal, while for glutamate we placed ROIs on the horizontal-cell dendrites immediately adjacent to a given cone, as previously (Yoshimatsu 656 et al., 2020b). To unequivocally identify UV-cones, responses to the 'red', 657 'green', 'blue' and 'UV'-flashes were always recorded (always 1 s flash, 1 s 658 darkness). Only cones that preferentially responded to the UV-LED were 659 kept for further analysis (Yoshimatsu et al., 2020a). 660
- Scaling and Denoising. We preprocessed the recorded and z-scored 662 fluorescence traces as follows (see Fig. S3 for a visualization): First, we 663 664 applied a linear baseline correction to the calcium traces to correct a linear 665 baseline decay. Then we rescaled the calcium and glutamate traces by z-666 scoring the traces with respect to the mean and standard deviation of the UV-bright stimulus intervals, resulting in a mean of zero and a standard 667 deviation of one in these intervals. By doing so we assumed that within these 668 periods calcium channels are closed and recorded activity is either due to 669 noise in the recording process or inherent channel/vesicle noise in the 670 synapse. With this normalization, we achieved a similar scaling for all traces 671 independent of the level of indicator expression. Based on these 672 preprocessed data (Fig. S3a,b second rows), the indices in Fig. 3c-j were 673 674 computed.

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For the model input, we further averaged each zone over the trials and then 675 applied a Butterworth filter of order three with a cut-off frequency of 5 Hz to 676 denoise the signals. Since the output of the model was in vesicles per 677 second, we finally shifted the processed glutamate data by its minimal value 678 such that it had only positive values. To use the calcium concentration as 679 input to the model, we additionally applied a Wiener deconvolution with the 680 kernel of the calcium indicator JRGeco1 to the data (assuming a SNR of 10 681 for f < 1Hz and a SNR of 1/20 for f > 1 Hz). The resulting pair of 682

calcium/glutamate data per zone (Fig. S3 last row, also Fig. 3a) was finally 683 used in the subsequent model. 684 685 Analysis of the SyGCaMP6f data from (Yoshimatsu et al., 2020b). 686 The z-scored (GCaMP6f) calcium data was smoothed with a sliding average 687 (window size of 100ms). Then two exponential functions (  $f(x) = c + a \cdot$ 688  $\exp(-\frac{1}{\tau} \cdot x)$ ) were fitted to the rise and decay periods. The time shift was 689 defined via the start of the calcium rise, more precisely as  $\min_{t} (Ca(t) > 3 \cdot$ 690 691 std(Ca)). 692 Data indices. Maximal activation, Sustain and Transience: To be less prone 693 to noise we used the 90<sup>th</sup>- and 50<sup>th</sup>-percentiles ( $p_i$ ) to calculate the maximal 694 activation, transience and sustain: 695  $\max = p_{90}(x_{[t_0,t_1]}),$ 696 sustain =  $p_{50}(x_{[t_2,t_3]})$ , 697 transience =  $\frac{\max - \text{sustain}}{\max}$ , 698 699 where x was the calcium/glutamate recording, respectively. The  $t_i$  were 700 chosen such that the first (for max) or the last (for sustained) second after 701 702 the onset of the dark period is included. Off-, On-Detection index (Ioff, Ion) and high frequency index (HFi): For the 703 detection indices we used a Gaussian noise stimulus (( $\mu, \sigma$ ) = (0.5,0.3)) 704 with 2 Hz and a length of 150 s. This stimulus was shown for 120 s before 705 the event to detect occurred with an amplitude of plus or minus four times 706 the standard deviation and a duration of 500 ms. The detection indices of the 707 simulation x were finally computed as 708 709  $I_{off}(x) = (\max(x_{[t_0,t_1]}) - \max(x_{[t_0,t_1]})) / \operatorname{std}(x_{[t_0,t_1]}),$ 710  $I_{on}(x) = abs(min(x_{[t_0,t_1]}) - mean(x_{[t_0,t_1]})) / std(x_{[t_0,t_1]})),$ 712 713 with  $t_0 = 60$  s and  $t_1 = 150$  s. 711 For the HFi we used uniformly distributed noise of 100 s at 20 Hz and 714 computed the discretized power spectrum p of the simulation x with Welch's 715 method, and defined the HFi on the by the standard deviation normalized 716 717 data x as  $HFi(x) = \sum_{i=1}^{n} p_i(x) \cdot f_i$ 718 with *n* such that  $f_n < 25$ Hz. 719

**Model.** We modelled the synaptic release by a cascade-like ribbon synapse model (Fig. 4a) with three vesicle pools (reserve pool *RP*, intermediate pool *IP* and readily releasable pool *RRP*) and changing rates which were dependent on the occupancy of the pools (Baden et al., 2014; Sterling and Matthews, 2005). In this model, the glutamate release e(t) was driven by the intracellular calcium Ca(t):

$$e(t) = e_{max} \cdot f(Ca(t)) \cdot \frac{RRP(t)}{RRP_{max}}$$

with

$$f(Ca) = \frac{1}{1 + \exp(-k \cdot (Ca - x_0))}.$$

As  $f(x_0) = 0.5$ , the parameter  $x_0$  specifies the operating point of the nonlinearity. It can be seen as an inverted baseline: the smaller  $x_0$  the less additional calcium is needed to trigger a vesicle release. If we assume a fixed calcium affinity for vesicle release, this implies an increased baseline level in the synapse.

The changing rates r(t) (between *RP* and *IP*) and i(t) (between the *IP* and *RRP*) were independent of calcium:

$$r(t) = r_{max} \cdot \left(1 - \frac{IP(t)}{IP_{max}}\right) \cdot \frac{RP(t)}{RP_{max}}$$
$$i(t) = i_{max} \cdot \left(1 - \frac{RRP(t)}{RRP_{max}}\right) \cdot \frac{IP(t)}{IP_{max}}$$

Additionally, the refilling d(t) of the reserve pool was modelled by a constant factor of the available exocytosed vesicles Exo(t):

$$d(t) = d_{max} \cdot Exo(t)$$

Therefore, the number of vesicles in the pools changed as:

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$$\frac{dRP(t)}{dt} = d(t) - r(t)$$

50 
$$\frac{dIP(t)}{dt} = r(t) - i(t)$$

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753  

$$\frac{dRRP(t)}{dt} = i(t) - e(t)$$

$$\frac{dExo(t)}{dt} = e(t) - d(t).$$

754In the original model in (Baden et al., 2014) there was also a non-linear755influence of calcium for i(t) in terms of  $\frac{Ca}{Ca+c}$ . But initial runs of the fitting756procedure resulted in  $c \approx 0$  which indicated no calcium dependency for the757changing rate *i* and we excluded this term in the presented model. Since the758endocytosis constant  $d_{max}$  and the size of the reserve pool RP did not affect

759 the output of the model, our fitting procedure was constrained to the remaining seven parameters: the changing rates (r<sub>max</sub>, *i<sub>max</sub>*, *e<sub>max</sub>*) the non-760 linearity parameters (k and  $x_0$ ), and the remaining pool sizes (IP and RRP). 761 We call these parameters  $\theta = (r_{max}, i_{max}, e_{max}, k, x_0, IP, RRP)$ . From the 762 equation above it follows that the model is scale invariant: a scaling of the 763 parameters (except the parameters for the non-linearity,  $x_0$  and k) results in 764 a scaled model output and thus only an arbitrary scale in vesicle units (v.u.) 765 766 to the experimental traces can be fitted.

- 767The described coupled ODE was solved with scipy's (version 1.5.1)768implementation of the Bogacki–Shampine method (Bogacki and Shampine,7691989), an explicit Runge-Kutta method of order 3 with adaptive step-sizes,770where the maximal step size was set to the step size of the (calcium) input771signal.
- 773Simplified Model. To reduce the parameter space to identify general rules774or ribbon tuning (Fig. 6d-f, Fig. 7 and online tool) we fixed k to 10.2 (equal to775the mean of the fitted parameters across zones) and additionally coupled the776maximal changing rates to the pool sizes as follows:

778	$r_{max} = 0.2 \cdot IP_{max}$
779	$i_{max} = 0.4 \cdot IP_{max}$
780	$e_{max} = \tilde{e}_{max} \cdot RRP_{max}$

781 where  $\tilde{e}_{max}$  can take values between 0 and 1.

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For the general rules of ribbon tuning (Fig. 6d-f, Fig. 7 and online tool) the calcium concentration evoked by a light stimulus s(t) is simulated as following:

 $Ca(t) = \kappa_2 * \exp(\kappa_1 * s(t))$ 

- 786where  $\kappa_1$  is a biphasic kernel from (Baden et al., 2014) and  $\kappa_2$  is a double787exponential kernel with fixed time constant  $\tau_{rise}$  (30 ms) and variable decay788parameter  $\tau_{decay}$ .
- Parameter Inference. For the inference of the model parameters, we
  iterated the following steps over several rounds: First we draw samples from
  a prior, evaluated the model and extracted summary statistics on which the
  loss was computed. Based on these loss values and the sampled
  parameters a mixture of density network (MDN) was trained and finally
  evaluated to get the posterior of this round which was used as the prior for
  the next round (Fig. 4b).
- 796 Summary statistics and relevant loss
- 797A key ingredient in simulation-based Bayesian inference is to define problem798and domain specific summary statistics to project the data to a low799dimensional feature space. For our setting, we identified the following800fourteen features x; baseline during adaption as well as period of UV-bright

801stimulus, mean during UV-dark period as well as mean of the maximal802release rates in this periods, maximal and minimal (as maximal value and80325<sup>th</sup> percentile) amplitude during the first flash, maximal and minimal (as804maximal value and 25<sup>th</sup> percentile) amplitude of the second peak, total805number of released vesicles during first, second and the last activation.

- 806To pay special attention to the decay after the initial UV-dark flash, which we807found to be informative about the different pool sizes of the ribbon, we fitted808an exponential decay to this period. We used the inferred time constant T809and the evaluation of the exponential function at the time point before the810next light onset as additional features along with an extra penalty if an811exponential rise instead of a decay was fitted.
- 812 To calculate the relevant loss R of a simulated trace e we normalized the 813 features in each component with the mean and standard deviation of the 814 recorded traces. We then took a weighted mean squared error of this 815 normalized summary vector x to the normalized summary vector  $x_0$  of the 816 recorded trace as the relevant loss:

$$R(e) = \frac{1}{14} \sum_{i=1}^{14} w_i \left( x_{0,i} - x_i \right)^2.$$

818The weights  $w_i$  were not systematically optimized, but within reasonable819ranges the results were relatively insensitive to the exact values. We chose820 $w = (0.5, 0.5, 5, 1, 1, 1, 1, 1, 1, 2, 1, 1, (0.01 for decay and <math>10 \cdot (1 + ceil(\tau))$  for rise))821for the features in the order mentioned as above, where the last value is the822extra penalty for the exponential rise instead of a decay.

## 823 Prior distribution and parameter normalization

- 824 The modes of the prior were chosen as  $(r, i, e, k, x_0, IP_{max}, RRP_{max}) = (2.5, RP_{max})$ 2.5, 10, 14, 0.5,  $\approx$ 13.8,  $\approx$ 4.0), but as the model was scale invariant, the 825 826 absolute values are uninformative and only the relative values are of interest. For technical reasons we normalized the prior distributions such that the 827 means of the uncorrelated multivariate normal distribution were 0.5 and the 828 829 standard deviations were 0.2 in each dimension. For the IP and RRP pool 830 sizes, we additionally exponentially scaled the sampled parameters, such 831 that no negative values could occur.
  - Parameter Inference

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We applied the Sequential Neural Posterior Estimation method described in 834 835 (Lueckmann et al., 2017) (code available at called https://github.com/mackelab/delfi) (also SNPE-B) with 836 some modifications which were also applied in (Oesterle et al., 2020; Yoshimatsu 837 838 et al., 2020a). In brief, SNPE-B draws over several rounds samples  $\{\theta_i\}_{i \in I}$ from a prior  $\tilde{p}(\theta)$  and evaluates the model for these parameters. For each 839 evaluation  $e_i$  the relevant loss function  $x_i = R(e_i)$  is computed and a mixture 840 841 density network (MDN)  $q_{\phi}(\theta, x)$  is trained on the data pairs  $\{(\theta_i, x_i)\}_{i \in I}$ . The posterior  $p(\theta|x_0)$  is then calculated as  $q_{\phi}(\theta|x = x_0)$  and used as a new 842 prior  $\tilde{p}(\theta)$  in the next sampling round. Instead of  $x_0 = 0$  we calculated the 843

new prior  $\tilde{p}(\theta)$  in round *n* as  $q_{\phi}(\theta | \mathbf{x} = \beta_n)$  where  $\beta_n$  is the 0.1<sup>th</sup> percentile  $\rho$ 844 of the relevant loss function of all samples. It turned out that this is an efficient 845 way to get a more stable behavior of the MDN since it has not to extrapolate 846 to unreached loss values but is converging nevertheless. This evaluation of 847 848  $q_{\phi}(\theta|x = \beta_n)$  at  $\beta_n = \rho$  can be seen as the posterior over the parameters for the "best-possible" model evaluations. Testing for different percentiles in a 849 850 reasonable range did not change the results. We took the same approach 851 for setting an adaptive bandwidth for the kernel. As an additional post-hoc 852 verification of the posteriors, we took as final posterior distributions the posterior of the round with the smallest median loss of its samples ("early 853 stopping"). 854 855

Technical details

We ran the inference algorithm over 5 rounds, with 300.000 samples per round. We chose three Gaussian components for the mixture of Gaussian distribution and a MDN with two hidden layers with 120 nodes each. In each round the network was trained for 800 epochs with a minibatch size of 1000. To let the MDN focus on regions of low relevant loss values, we used a combined half-uniform-half-Gaussian kernel which was constant up to the pseudo observation  $\beta_n$  and decayed then as a half Gaussian. The scale of this half-Gaussian part of the kernel was in each round chosen as the 25<sup>th</sup>percentile of the relevant loss function.

867Sensitivity Analysis. Sensitivity analysis was performed using uncertainpy868(Tennøe et al., 2018) by using the GaussianMixture probability distribution869class of chaospy (Feinberg and Langtangen, 2015). Since model evaluations870are computationally cheap and could run in parallel, we took the (quasi-)871Monte Carlo Method with 10e5 samples, resulting in 450.000 simulations.872With uncertainpy we calculated the first order Sobol sensitivity index S<sub>i</sub> which873is defined as

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 $S_i = \frac{\mathbb{V}[\mathbb{E}[Y|\theta_i]]}{\mathbb{V}[Y]},$ 

where  $\mathbb{E}$  and  $\mathbb{V}$  are the expected value and the variance, respectively.  $S_i$ 875 measures the direct effect each parameter has on the variance of the model 876 output (Saltelli et al., 2008; Tennøe et al., 2018). It tells us the expected 877 reduction in relative variance if we fix parameter  $\theta_i$ . The sum of the first-order 878 Sobol indices cannot exceed one and is equal to one if no interactions are 879 present (Glen and Isaacs, 2012; Tennøe et al., 2018). We also calculated 880 881 the total-order indices which gives the sensitivity due to interactions of the 882 parameters. But since already the sum of the first-order indices is almost one 883 for our model, the total order indices look quite similar and we omit their analysis. 884

885It is important to remark, that the Sobol indices are not a way to show how886'important' parameters are. A parameter which is overbearingly influencing887the model would in the limit of the posterior estimation accumulate the mass888on one single point. But this means that for samples from the posterior this

889 parameter is not responsible for any variance of the model output and thus its Sobol index would be zero. The Sobol indices are therefore the result of 890 a complex interplay of the model and the posterior estimation, but especially 891 its temporal changes give us insight into the time dependent influence on the 892 model output. However, the posteriors are an adequate probability 893 distribution to calculate the Sobol indices, as they provide all parameter 894 895 combinations which are in agreement with the experimental data and are 896 more expressive as commonly used uniform distributions, where each 897 marginal is simply defined as the mean ±10% of the fitted parameters.

- 898Linear baseline model. To evaluate the performance of the biophysical899model, we compared it to a simple linear model. For this, we performed a900regularized least square regression (ridge regression using scikit-learn,901version 0.23.1, https://scikit-learn.org) to fit the calcium to glutamate902response. In lack of a diverse enough dataset, we set the regularization903coefficient alpha to 0.1 and allowed the model to include the data from the904past 0.5 second to predict the next time point.
- 905 Statistical Analysis
- 906 Vesicle Densities
- We used Generalized Additive Models (GAMs) for the comparison of the 907 vesicle densities (Fig. 1g, Fig. S1). GAMs are an extension to generalized 908 909 linear models by allowing linear predictors that depend on smooth functions 910 of the underlying variables (Wood, 2017). We used the mgcv-package 911 (version 1.8-33) in R on an Windows 10 workstation with default parameters, if not specified differently below. We modelled the dependence of the vesicle 912 density as a smooth term dependent on the distance with 100 degrees of 913 freedom and grouped by "zone". We further used "zone" as additional 914 predictive variable. The model explained ~65% of the deviance. Statistical 915 significance for differences between the dependence on the vesicle density 916 917 in the different retinal regions were obtained using the *plot\_diff* function of the *itsaduq*-package for R (version 2.4) with a 95% confidence level. 918
- 919 Hypothesis Testing
- 920For comparisons between regional properties (ribbon geometry (Fig. 1) and921functional indices (Fig. 3)) we used two-sided shuffling tests with Bonferroni922correction. Sample sizes and significant levels are stated in the figure923captions.

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